# PARHAM, et al. Polyulistide Croding DIES! ATUMAN RECEPTOR PROTEINS; RELATED REAGENTS AND METHODS

This filing is a conversion of U.S. Provisional Patent Application 60/077,329, filed March 9, 1998, which is incorporated herein by reference, to a U.S. Utility Patent Application.

## FIELD OF THE INVENTION

The present invention relates to compositions and
methods for affecting mammalian physiology, including
morphogenesis or immune system function. In particular, it
provides nucleic acids, proteins, and antibodies which
regulate development and/or the immune system. Diagnostic
and therapeutic uses of these materials are also disclosed.

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## BACKGROUND OF THE INVENTION

Recombinant DNA technology refers generally to techniques of integrating genetic information from a donor source into vectors for subsequent processing, such as through introduction into a host, whereby the transferred genetic information is copied and/or expressed in the new environment. Commonly, the genetic information exists in the form of complementary DNA (cDNA) derived from messenger RNA (mRNA) coding for a desired protein product. The carrier is frequently a plasmid having the capacity to incorporate cDNA for later replication in a host and, in some cases, actually to control expression of the cDNA and thereby direct synthesis of the encoded product in the host. See, e.g., Sambrook, et al. (1989) Molecular Cloning: A Laboratory Manual, (2d ed.), vols. 1-3, CSH Press, NY.

For some time, it has been known that the mammalian immune response is based on a series of complex cellular interactions, called the "immune network". Recent research has provided new insights into the inner workings of this network. While it remains clear that much of the immune response does, in fact, revolve around the network-like

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interactions of lymphocytes, macrophages, granulocytes, and other cells, immunologists now generally hold the opinion that soluble proteins, known as lymphokines, cytokines, or monokines, play critical roles in controlling these cellular interactions. The interferons are generally considered to be members of the cytokine family. Thus, there is considerable interest in the isolation, characterization, and mechanisms of action of cell modulatory factors, an understanding of which will lead to significant advancements in the diagnosis and therapy of numerous medical abnormalities, e.g., immune system disorders.

Lymphokines apparently mediate cellular activities in a variety of ways. See, e.g., Paul (ed. 1996) Fundamental Immunology 3d ed., Raven Press, New York; and Thomson (ed. 1994) The Cytokine Handbook 2d ed., Academic Press, San Diego. They have been shown to support the proliferation, growth, and/or differentiation of pluripotential hematopoietic stem cells into vast numbers of progenitors comprising diverse cellular lineages which make up a complex immune system. Proper and balanced interactions between the cellular components are necessary for a healthy immune response. The different cellular lineages often respond in a different manner when lymphokines are administered in conjunction with other agents.

Cell lineages especially important to the immune response include two classes of lymphocytes: B-cells, which can produce and secrete immunoglobulins (proteins with the capability of recognizing and binding to foreign matter to effect its removal), and T-cells of various subsets that secrete lymphokines and induce or suppress the B-cells and various other cells (including other T-cells) making up the immune network. These lymphocytes interact with many other cell types.

One means to modulate the effect of a cytokine upon binding to its receptor, and therefore potentially useful in treating inappropriate immune responses, e.g., autoimmune,

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inflammation, sepsis, and cancer situations, is to inhibit the receptor signal transduction. Unfortunately, finding reagents capable of serving as an antagonist or agonist has been severely hampered by the failure to fully identify all of the components within the signaling systems. In order to characterize the structural properties of a cytokine receptor in greater detail and to understand the mechanism of action at the molecular level, purified receptor will be very useful. The receptors provided herein, by comparison to other receptors or by combining structural components, will provide further understanding of signal transduction induced by ligand binding.

The isolated receptor gene should provide means to generate an economical source of the receptor, allow expression of more receptors on a cell leading to increased assay sensitivity, promote characterization of various receptor subtypes and variants, and allow correlation of activity with receptor structures. Moreover, fragments of the receptor may be useful as agonists or antagonists of ligand binding. See, e.g., Harada, et al. (1992) J. Biol. Chem. 267:22752-22758. Often, there are at least two critical subunits in the functional receptor. See, e.g., Gonda and D'Andrea (1997) Blood 89:355-369; Presky, et al. (1996) Proc. Nat'l Acad. Sci. USA 93:14002-14007; Drachman and Kaushansky (1995) Curr. Opin. Hematol. 2:22-28; Theze (1994) Eur. Cytokine Netw. 5:353-368; and Lemmon and Schlessinger (1994) Trends Biochem. Sci. 19:459-463.

From the foregoing, it is evident that the discovery and development of new soluble proteins and their receptors, including ones similar to lymphokines, should contribute to new therapies for a wide range of degenerative or abnormal conditions which directly or indirectly involve development, differentiation, or function, e.g., of the immune system and/or hematopoietic cells. In particular, the discovery and understanding of novel receptors for lymphokine-like molecules which enhance or potentiate the beneficial activities of other

lymphokines would be highly advantageous. The present invention provides new receptors for ligands exhibiting similarity to cytokine like compositions and related compounds, and methods for their use.

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## SUMMARY OF THE INVENTION

The present invention is directed to novel receptors related to cytokine receptors, e.g., primate or rodent, cytokine receptor like molecular structures, designated DNAX Interferon-like Receptor Subunits (DIRS), and their biological activities. In particular, it provides description of two different subunits, designated DIRS1 and DIRS2. It includes nucleic acids coding for the polypeptides themselves and methods for their production and use. The nucleic acids of the invention are characterized, in part, by their homology to cloned complementary DNA (cDNA) sequences enclosed herein.

The present invention provides, in polypeptide embodiments: a substantially pure or recombinant DIRS1 polypeptide comprising at least three distinct nonoverlapping segments of at least four amino acids identical to segments of SEQ ID NO: 2; a substantially pure or recombinant DIRS1 polypeptide comprising at least two distinct nonoverlapping segments of at least five amino acids identical to segments of SEQ ID NO: 2; a natural sequence DIRS1 comprising mature SEQ ID NO: 2; a fusion polypeptide comprising DIRS1 sequence; a substantially pure or recombinant DIRS2 polypeptide comprising at least three distinct nonoverlapping segments of at least ten amino acids identical to segments of SEQ ID NO: 4; a substantially pure or recombinant DIRS2 polypeptide comprising at least two distinct nonoverlapping segments of at least eleven amino acids identical to segments of SEQ ID NO: 4; a natural sequence DIRS2 comprising SEQ ID NO: 4; or a fusion polypeptide comprising DIRS2 sequence. Preferred embodiments include, e.g., the substantially pure or isolated antigenic: DIRS1 polypeptide, wherein the distinct nonoverlapping segments of identity: include one of at least eight amino acids; include one of at least four amino acids and a second of at least five amino acids; include at least three segments of at least four, five, and six amino

acids, or include one of at least twelve amino acids; or

DIRS2 polypeptide, wherein the distinct nonoverlapping segments of identity: include one of at least thirteen amino acids; include one of at least eleven amino acids and a second of at least thirteen amino acids; include at least three segments of at least ten, eleven, and twelve amino acids; or include one of at least twenty-five amino acids. Other embodiments include compositions where: the DIRS1 polypeptide: comprises a mature sequence of Table 1; is an unglycosylated form of DIRS1; is from a primate, such as a human; comprises at least seventeen amino acids of SEQ ID 10 NO: 2; exhibits at least four nonoverlapping segments of at least seven amino acids of SEQ ID NO: 2; is a natural allelic variant of DIRS1; has a length at least about 30 amino acids; exhibits at least two non-overlapping epitopes which are specific for a primate DIRS1; is glycosylated; 15 has a molecular weight of at least 30 kD with natural glycosylation; is a synthetic polypeptide; is attached to a solid substrate; is conjugated to another chemical moiety; is a 5-fold or less substitution from natural sequence; or is a deletion or insertion variant from a natural sequence; 20 or the DIRS2 polypeptide: comprises a mature sequence of Table 2; is an unglycosylated form of DIRS2; or is from a primate, such as a human; comprises at thirty-five amino acids of SEQ ID NO: 4; exhibits at least four 25 nonoverlapping segments of at least twelve amino acids of SEQ ID NO: 4; is a natural allelic variant of DIRS2; has a length at least about 30 amino acids; exhibits at least two non-overlapping epitopes which are specific for a primate DIRS2; is glycosylated; has a molecular weight of at least 30 kD with natural glycosylation; is a synthetic 30 polypeptide; is attached to a solid substrate; is conjugated to another chemical moiety; is a 5-fold or less substitution from natural sequence; or is a deletion or insertion variant from a natural sequence. Various 35 combination compositions include those comprising: a substantially pure DIRS1 and another Interferon Receptor

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family member; a substantially pure DIRS2 and another

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Interferon Receptor family member; a sterile DIRS1 polypeptide; a sterile DIRS2 polypeptide; the DIRS1 polypeptide and a carrier, wherein the carrier is: an aqueous compound, including water, saline, and/or buffer; and/or formulated for oral, rectal, nasal, topical, or parenteral administration; or the DIRS2 polypeptide and a carrier, wherein the carrier is: an aqueous compound, including water, saline, and/or buffer; and/or formulated for oral, rectal, nasal, topical, or parenteral administration.

Fusion polypeptide embodiments include those comprising: mature protein sequence of Table 1; mature protein sequence of Table 2; a detection or purification tag, including a FLAG, His6, or Ig sequence; or sequence of another interferon receptor protein. Kit embodiments are provided, e.g., a kit comprising such a polypeptide, and: a compartment comprising the protein or polypeptide; or instructions for use or disposal of reagents in the kit.

The invention also provides a binding compound comprising an antigen binding site from an antibody, which specifically binds to a: natural DIRS1 polypeptide, wherein: the binding compound is in a container; the DIRS1 polypeptide is from a human; the binding compound is an Fv, Fab, or Fab2 fragment; the binding compound is conjugated to another chemical moiety; or the antibody: is raised against a peptide sequence of a mature polypeptide of Table 1; is raised against a mature DIRS1; is raised to a purified human DIRS1; is immunoselected; is a polyclonal antibody; binds to a denatured DIRS1; exhibits a Kd to antigen of at least 30  $\mu\text{M}$ ; is attached to a solid substrate, including a bead or plastic membrane; is in a sterile composition; or is detectably labeled, including a radioactive or fluorescent label; or a natural DIRS2 polypeptide, wherein: the binding compound is in a container; the DIRS2 protein is from a human; the binding compound is an Fv, Fab, or Fab2 fragment; the binding compound is conjugated to another chemical moiety; or the

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antibody: is raised against a peptide sequence of a mature polypeptide of Table 2; is raised against a mature DIRS2; is raised to a purified human DIRS2; is immunoselected; is a polyclonal antibody; binds to a denatured DIRS2; exhibits a Kd to antigen of at least 30 µM; is attached to a solid substrate, including a bead or plastic membrane; is in a sterile composition; or is detectably labeled, including a radioactive or fluorescent label. Kit embodiments include, e.g., those comprising the binding compound, and: a compartment comprising the binding compound; or instructions for use or disposal of reagents in the kit.

Various methods are provided, e.g., of producing an antigen:antibody complex, comprising contacting under appropriate conditions: a primate DIRS1 polypeptide with a described antibody; or a primate DIRS2 polypeptide with a described antibody; thereby allowing the complex to form. In certain situations, the method is used wherein: the complex is purified from other interferon receptors; the complex is purified from other antibody; the contacting is with a sample comprising an interferon; the contacting allows quantitative detection of the antibody; or the contacting allows quantitative detection of the antibody; or the contacting allows quantitative detection of the antibody.

Other compositions comprise: a sterile binding
compound as described, or the described binding compound
and a carrier, wherein the carrier is: an aqueous compound,
including water, saline, and/or buffer; and/or formulated
for oral, rectal, nasal, topical, or parenteral
administration.

Nucleic acid embodiments include, e.g., an isolated or recombinant nucleic acid encoding the: described DIRS1 polypeptide, wherein the: DIRS1 is from a human; or the nucleic acid: encodes an antigenic peptide sequence of Table 1; encodes a plurality of antigenic peptide sequences of Table 1; exhibits identity over at least thirteen nucleotides to a natural cDNA encoding the segment; is an expression vector; further comprises an origin of

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replication; is from a natural source; comprises a detectable label; comprises synthetic nucleotide sequence; is less than 6 kb, preferably less than 3 kb; is from a primate; comprises a natural full length coding sequence; is a hybridization probe for a gene encoding the DIRS1; or is a PCR primer, PCR product, or mutagenesis primer; or the described DIRS2 polypeptide, wherein the: DIRS2 is from a human; or the nucleic acid: encodes an antigenic peptide sequence of Table 2; encodes a plurality of antigenic peptide sequences of Table 2; exhibits identity over at least 30 nucleotides to a natural cDNA encoding the segment; is an expression vector; further comprises an

origin of replication; is from a natural source; comprises

sequence; is less than 6 kb, preferably less than 3 kb; is 15 from a primate; comprises a natural full length coding sequence; is a hybridization probe for a gene encoding the DIRS2; or is a PCR primer, PCR product, or mutagenesis

a detectable label; comprises synthetic nucleotide

primer.

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20 The invention further provides a cell or tissue comprising the described recombinant nucleic acid. Certain embodiments include wherein the cell is: a prokaryotic cell; a eukaryotic cell; a bacterial cell; a yeast cell; an insect cell; a mammalian cell; a mouse cell; a primate cell; or a human cell. Kits are also provided, e.g., the 25 described nucleic acid and: a compartment comprising the nucleic acid; a compartment further comprising a primate DIRS1 polypeptide; a compartment further comprising a primate DIRS2 polypeptide; or instructions for use or 30 disposal of reagents in the kit.

In other embodiments, the invention provides a nucleic acid which: hybridizes under wash conditions of 30 minutes at 30° C and less than 2M salt to the coding portion of SEQ ID NO: 1; hybridizes under wash conditions of 30 minutes at  $30^{\circ}$  C and less than 2M salt to the coding portion of SEQ ID NO: 3; exhibits identity over a stretch of at least about 30 nucleotides to a primate DIRS1 sequence; or exhibits

identity over a stretch of at least about 30 nucleotides to a primate DIRS2 sequence. Preferred embodiments include those nucleic acids wherein: the wash conditions are at 45° C and/or 500 mM salt; or the stretch is at least 55 nucleotides. Other embodiments include those nucleic acids wherein: the wash conditions are at 55° C and/or 150 mM salt; or the stretch is at least 75 nucleotides.

The invention further provides a method of modulating physiology or development of a cell or tissue culture cells comprising contacting the cell with an agonist or antagonist of a mammalian DIRS1 or DIRS2. The method may involve where the cell is transformed with a nucleic acid encoding a DIRS1 or DIRS2 and another cytokine receptor subunit.

## DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

### OUTLINE

I	G	en	e	r	a	1

- 5 II. Activities
  - III. Nucleic acids
    - A. encoding fragments, sequence, probes
    - B. mutations, chimeras, fusions
    - C. making nucleic acids
- 10 D. vectors, cells comprising
  - IV. Proteins, Peptides
    - A. fragments, sequence, immunogens, antigens
    - B. muteins
    - C. agonists/antagonists, functional equivalents
- D. making proteins
  - V. Making nucleic acids, proteins
    - A. synthetic
    - B. recombinant
    - C. natural sources
- 20 VI. Antibodies
  - A. polyclonals
  - B. monoclonal
  - C. fragments; Kd
  - D. anti-idiotypic antibodies
- 25 E. hybridoma cell lines
  - VII. Kits and Methods to quantify DIRS
    - A. ELISA
    - B. assay mRNA encoding
    - C. qualitative/quantitative
- 30 D. kits
  - VIII. Therapeutic compositions, methods
    - A. combination compositions
    - B. unit dose
    - C. administration
- 35 IX. Screening
  - X. Ligands

## I. General

The present invention provides the amino acid
sequences and DNA sequences of mammalian, herein primate,
interferon receptor-like subunit molecules, these ones
designated DNAX Interferon Receptor family Subunit 1
(DIRS1) and DNAX Interferon Receptor family Subunit 2,
having particular defined properties, both structural and

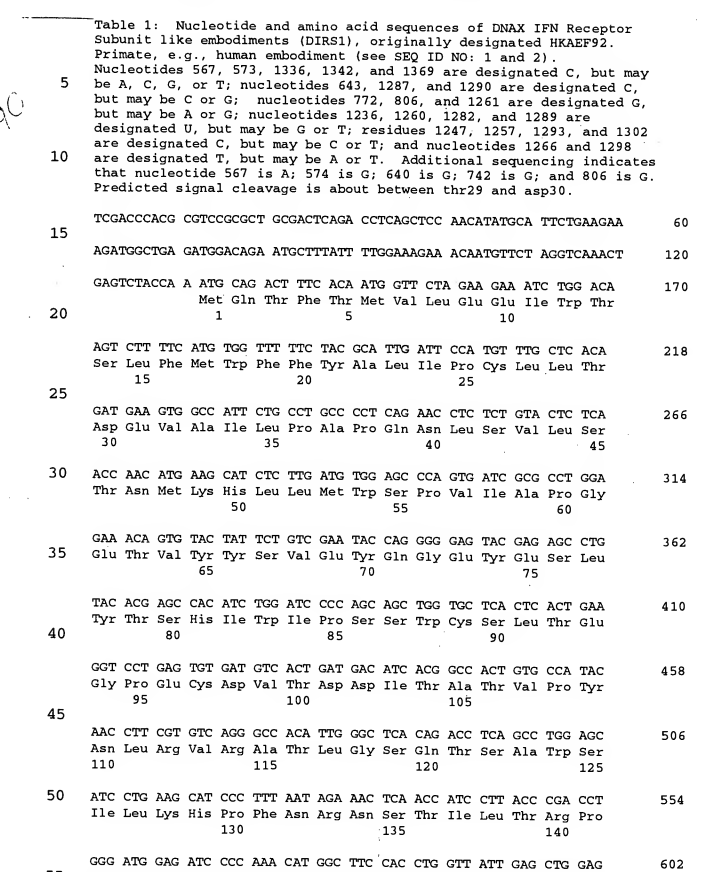
biological. Various cDNAs encoding these molecules were obtained from primate, e.g., human, cDNA sequence libraries. Other primate or other mammalian counterparts

would also be desired. Descriptions, methods, and manipulations directed to DIRS1 may be applied, as appropriate, to DIRS2.

Some of the standard methods applicable are described or referenced, e.g., in Maniatis, et al. (1982) Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor Press; Sambrook, et al. (1989) Molecular Cloning: A Laboratory Manual, (2d ed.), vols. 1-3, CSH Press, NY; Ausubel, et al., Biology, Greene Publishing Associates, Brooklyn, NY; or Ausubel, et al. (1987 and periodic supplements) Current Protocols in Molecular Biology, Greene/Wiley, New York; each of which is incorporated herein by reference.

A partial nucleotide (SEQ ID NO: 1) and corresponding 15 amino acid sequence (SEQ ID NO: 2) of a human DIRS1 coding segment is shown in Table 1. Partial human DIRS2 sequence is provided (SEQ ID NO: 3 and 4).

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Gly Met Glu Ile Pro Lys His Gly Phe His Leu Val Ile Glu Leu Glu

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	GAC Asp	CTG Leu	GGG Gly 160	CCC Pro	CAG Gln	TTT Phe	GAG Glu	TTC Phe 165	CTT Leu	GTG Val	GCC Ala	TAC Tyr	TGG Trp 170	ACG Thr	AGG Arg	GAG Glu		650
5	CCT Pro	GGT Gly 175	GCC Ala	GAG Glu	GAA Glu	CAT His	GTC Val 180	AAA Lys	ATG Met	GTG Val	AGG Arg	AGT Ser 185	GGG Gly	GGT Gly	ATT Ile	CCA Pro		698
10				GAA Glu														746
15	CAG Gln	ACA Thr	TTC Phe	GTG Val	AAG Lys 210	GCC Ala	ATT Ile	GGG Gly	AGG Arg	TAC Tyr 215	AGC Ser	GCC Ala	TTC Phe	AGC Ser	CAG Gln 220	ACA Thr		794
20	GAA Glu	TGT Cys	GTG Val	GAG Glu 225	GTG Val	CAA Gln	GGA Gly	GAG Glu	GCC Ala 230	ATT Ile	CCC Pro	CTG Leu	GTA Val	CTG Leu 235	GCC Ala	CTG Leu		842
				GTT Val														890
25	GTC Val	TGG Trp 255	AAA Lys	ATG Met	GGC Gly	CGG Arg	CTG Leu 260	CTC Leu	CAG Gln	TAC Tyr	TCC Ser	TGT Cys 265	TGC Cys	CCC Pro	GTG Val	GTG Val		938
30	GTC Val 270	CTC Leu	CCA Pro	GAC Asp	ACC Thr	TTG Leu 275	AAA Lys	ATA Ile	ACC Thr	AAT Asn	TCA Ser 280	CCC Pro	CAG Gln	AAG Lys	TTA Leu	ATC Ile 285		986
35	AGC Ser	TGC Cys	AGA Arg	AGG Arg	GAG Glu 290	GAG Glu	GTG Val	GAT Asp	GCC Ala	TGT Cys 295	GCC Ala	ACG Thr	GCT Ala	GTG Val	ATG Met 300	TCT Ser	1	034
40	CCT Pro	GAG Glu	GAA Glu	CTC Leu 305	CTC Leu	AGG Arg	GCC Ala	TGG Trp	ATC Ile 310	TCA Ser	TAGG	TTTC	ece è	AAGG	GCCC	:A	1	084
- •	GGTG	AAGO	CCG A	AGAAC	CTGG	T CI	GCAT	GACA	TGG	AAAC	CAT	GAGG	GGAC	AA G	TTGT	GTTT	rc 1	144
	TGTT	TTCC	CC C	CACGG	ACAA	'G GG	ATGA	GAGĄ	AGI	'AGGA	AGA	GCCI	GTTG	TC T	ACAA	GTCT	'A 1	204
45	GAAG	CAAC	CA T	CAGA	.GGCA	G GG	TGGT	TTGI	CTA	ACAG	AAC	AACT	GACI	'GA G	GCTA	TGGG	G 1	264
	GTTG	TGAC	CT C	TAGA	CTTT.	'G GG	CTTC	CACI	TGC	TTGG	CTG	AGCA	ACCC	TG G	GAAA	AGTG	A 1	324
	CTTC	ATCC	CT I	CGGT	CCCA	A GT	TTTC	TCAT	CTG	TAAT	'GGG	GGAT	CCCT	AC A	AAAC	TG	1	381

	5	Rec HOF	epto NY28	r Su (SE	buni Q ID	t li NO:	ke e 3 a	mbod nd 4	imen ).	ts ( Nucl	DIRS eoti	2), de 1	orig	inal esig	ly d nate	esig d C.	AX IF nated may		
100	J	C C	GG G rg V 1	TC G	AC C sp P	CA C	GC G rg V 5	TC C	GC C' rg L	TG G eu V	al S	CC C er P 10	CC T	GG C	TG A eu T	hr V	TG al 15		46
`	10	CCT Pro	TGG Trp	TTC Phe	CTG Leu	TCC Ser 20	Cys	TGG Trp	AAT Asn	GTT Val	ACC Thr 25	ATT Ile	GGG	CCT Pro	CCT Pro	GAG Glu 30	AGC Ser		94
	15	ATC Ile	TGG Trp	GTG Val	ACG Thr 35	CCG Pro	GGA Gly	GAA Glu	GCC Ala	TCC Ser 40	CTC Leu	ATC Ile	AȚC Ile	AGG Arg	TTC Phe 45	TCC Ser	TCT Ser		142
	20	CCC Pro	TTC Phe	GAC Asp 50	GTC Val	CCT Pro	CCC Pro	AAC Asn	CTG Leu 55	GGC Gly	TAT Tyr	TTC Phe	CAG Gln	TAC Tyr 60	TAT Tyr	GTC Val	CAT His		190
	25	TAC Tyr	TGG Trp 65	GAA Glu	AAG Lys	GCG Ala	GGA Gly	ATC Ile 70	CAA Gln	AAG Lys	GTT Val	AAA Lys	GGT Gly 75	CCT Pro	TTC Phe	AAG Lys	AGC Ser		238
		AAC Asn 80	TCC Ser	ATC	GTG Val	TTG Leu	GAT Asp 85	GGC Gly	TTG Leu	AGA Arg	CCC Pro	TTA Leu 90	AGA Arg	GAA Glu	TAC Tyr	TGT Cys	TTÄ Leu 95		286
	30	CAA Gln	GTG Val	AAG Lys	GCG Ala	CAT His 100	CTC Leu	TTT Phe	CGC Arg	ACA Thr	TCC Ser 105	TGC Cys	AAC Asn	ACC Thr	TCT Ser	AGG Arg 110	CCC Pro		334
	35	GGC	CGC Arg	TTA Leu	AGC Ser 115	AAC Asn	ATA Ile	ACT Thr	TGC Cys	TAC Tyr 120	GAA Glu	ACA Thr	ATG Met	ATG Met	GAT Asp 125	GCC Ala	ACT Thr		382
	40	ACG Thr	AAG Lys	CTT Leu 130	CAA Gln	CAA Gln	GTC Val	ATC Ile	CTC Leu 135	ATC Ile	GCC Ala	GTG Val	GGA Gly	GTC Val 140	TTT Phe	CTG Leu	TCG Ser	,	430
	45	CTG Leu	GCG Ala 145	GCG Ala	CTG Leu	GCG Ala	GGG Gly	GGC Gly 150	TGT Cys	TTC Phe	TTC Phe	CTG Leu	GTG Val 155	CTG Leu	AGA Arg	TAC Tyr	AAA Lys		478
		GGC Gly 160	CTG Leu	GTG Val	AAA Lys	TAC Tyr	TGG Trp 165	TTT Phe	CAC His	TCT Ser	CCG Pro	CCA Pro 170	AGC Ser	ATC Ile	CCA Pro	TCA Ser	CAA Gln 175	!	526
	50												ATC Ile					!	574

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				GCC TGG GAC TTG ( Ala Trp Asp Leu \ .	
<b>.</b>	GTT GCA TT Val Ala Ph 21	ne Pro Ala Ly	G GAG CAA GAA ( s Glu Gln Glu ) 215	GAT GTT CCC CAA A Asp Val Pro Gln S 220	AGC ACT TTG 670 Ser Thr Leu
10		AC TCT GGT GC sn Ser Gly Al		CTGTGG GGTAAGGGC1	CTGAGCCGAG 724
	GAAGCTGCTG	G ATGTCCATGT	CAGCACTTTA TGG	AATCCGG TCCTCCATT	TT TCCTGTCCCC 784
15	AAAAGGCCCG	TCAGTGCCTG	rgaagatgta acg	GGTCTCA TGGGGGCG	AC AAGCTTATTG 844
	ATTTTTTTCT	TCAAACTAAG	AGTTTTCTAA TCA	TACGCGT TTTTAGAA1	A ATTCTACAGA 904
20	TATGTCCCCG	AAAGATTAAG	ATTTCTCTTA AAC	ACTAAAA AGACATGTA	A TTATTTGTTA 964
20	GCAAATGGGC	GTCTGGCACG	CCTCTGACAC TTT	TTCGTCA GCAGCCAGG	SA CACGAGGTCC 1024
	CCTCCTTGAT	GAAGCCCCTC (	GGCAGACCA TGT	CACCTGT CCCAGCCTG	C CCCAAGAAGG 1084
25	GACATTAAGT	GGCCCTTCTT (	CATATCCAAA CAC	CTGGCTT GAAATGTG#	T TAGCCCTGTA 1144
	AATAGTTTCA	CAGAGATTAA (	GCCTTTTTTT CCC	CCAAGTT AGGAATAAA	A GACTATAATT 1204
20	AACTTTTTAA	AAAAAAAAA A	LAAA AAAAAAAA	AAAAAA	1244

	5	DR1 prot (SEQ the	e 3: Sequer is a primate ein sequence ID NO: 5), crf2-4 prote mics 16:366	e DIRS1 prof e; the IRβ : see Soh, ef ein (SEQ ID	tein sequend is the human t al. (1994)	ce; DR2 is a n IFN-γ rece n <u>Cell</u> 76:79	a primate Di eptor beta s 93-802; and	IRS2 subunit CRF is
0120	10	DR2 DR1	MQTFTMVLEE	IWTSLFMWFF	YALIPCLLTD FAAAAAAPPD	EVAILPAPQN PLSQLPAPQH	LSVLSTNMKH PKIRLYNAEO	LLMWSPVIAP VLSWEPVALS
	15	DR2 DR1 IRβ crf	GETVYYSVEY NSTRPVVYRV KGNLTFTAQY	QGEYESLY QFKYTDSKWF LSYR	TSHIWIPSSW TADIMSIGVN	CSLTEGPECD CTQITATECD	FTAASPSAGF	VPYNLRV PMDFNVTLRL
	20	DR2 DR1 IRβ crf	RATLGSQTSA RAELGALHSA	WNVTIGPPES WSILK-HPFN WVTMPWFQHY WVNIT-FCPV	RNSTILTRPG RNVTVGPPEN	MEIXKXGFHL IEVTPGEGSL	VIELEDL IIRFSSPFDI	GPQADTS
	30	DR2 DR1 IRβ	FEFLVA -TAFFCYYVH	KVKGPFKSNS YWXREPGAEE YWEKGGIQ YWKNGTDE	HVKMVRSGGI QVKGPFRSNS	PVHLETMEPG -ISLDNLKPS	AAYCVKAQT- RVYCLQVQAQ	-FVKAIGX LLWNKSNIFR
	35	DR2 DR1 IRβ crf	YSAFSQTECV VGHLSNISCY	QQVILIAVGV EVQG-EAIPL ETMADASTEL QTTHDETVPS	VLALFAFVG- QQVILISVGT	-FMLILVVVP FSLLSVLAGA	LFVWKMGR CFFLVLKYRG	LLQYSCCPVV LIKYWFHTPP
	40	DR2 DR1 IRβ crf	VLPDTLKITN SIPLQIEEYL	LDKDTSPTDD S-P-QKLISC KDPTQPILEA GHPHHNTLLF	RREEVD LDKDSSPKDD	ACATAVMS VWDSVSIISF	PEE PEKEQE	
	45	DR2 DR1 IRβ crf	SGAVC -LLRAWIS DVLQTL DSCSLGTPPG	QGPQS				

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Table 3 shows comparison of the available sequences of primate embodiments of DIRS1, DIRS2, and two related interferon receptor family members. Both of the new DIRS appear to exhibit sequence similarity to beta interferon receptor subunits.

Structural features of the human DIRS1, and similarly for the other receptors as aligned in Table 3, include characteristic transmembrane segments of the IR $\beta$  and crf from 261-273, and correspond to: from about vall to pro133; fibronectin domains corresponding to the DIRS1 sequence from about gly134 to pro232, gly233 to gly306, and pro307 to lys403; a transmembrane segment from about val404 to gly427; and an intracellular domain from about arg428 to the carboxy terminus. Of particular interest is the WGEWS motif corresponding to residues trp104 to ser108.

As used herein, the term DIRS1 shall be used to describe a protein comprising a protein or peptide segment having or sharing the amino acid sequence shown in Table 1, or a substantial fragment thereof. The invention also includes a protein variation of the respective DIRS1 allele whose sequence is provided, e.g., a mutein or soluble extracellular construct. Typically, such agonists or antagonists will exhibit less than about 10% sequence differences, and thus will often have between 1- and 11fold substitutions, e.g., 2-, 3-, 5-, 7-fold, and others. It also encompasses allelic and other variants, e.g.,

natural polymorphic, of the protein described. Typically, it will bind to its corresponding biological ligand, perhaps in a dimerized state with an alpha receptor subunit, with high affinity, e.g., at least about 100 nM, usually better than about 30 nM, preferably better than about 10 nM, and more preferably at better than about 3 nM. The term shall also be used herein to refer to related naturally occurring forms, e.g., alleles, polymorphic variants, and metabolic variants of the mammalian protein.

This invention also encompasses proteins or peptides having substantial amino acid sequence identity with the amino acid sequence in Table 1. It will include sequence

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variants with relatively few substitutions, e.g., preferably less than about 3-5. Other embodiments include forms in association with an alpha subunit, e.g., a DSRS1, and/or with ligand, e.g., DIL-30.

A substantial polypeptide "fragment", or "segment", is a stretch of amino acid residues of at least about 8 amino acids, generally at least 10 amino acids, more generally at least 12 amino acids, often at least 14 amino acids, more often at least 16 amino acids, typically at least 18 amino acids, more typically at least 20 amino acids, usually at least 22 amino acids, more usually at least 24 amino acids, preferably at least 26 amino acids, more preferably at least 28 amino acids, and, in particularly preferred embodiments, at least about 30 or more amino acids, e.g., 35, 40, 50, 70, 90, 110, etc. Specific ends may be at all possible or appropriate combinations, or at proline residues. Sequences of segments of different proteins can be compared to one another over appropriate length stretches.

The invention provides polypeptides exhibiting a plurality of distinct, e.g., nonoverlapping, segments of the specified length. Typically, the plurality will be at least two, more usually at least three, and preferably 5, 7, or even more. While the length minima are provided, longer lengths, of various sizes, may be appropriate, e.g., one of length 7, and two of length 12.

Amino acid sequence homology, or sequence identity, is determined by optimizing residue matches. In some comparisons, gaps may be introduces, as required. See, e.g., Needleham, et al. (1970) <u>J. Mol. Biol.</u> 48:443-453; Sankoff, et al., (1983) chapter one in <u>Time Warps</u>, <u>String Edits</u>, and <u>Macromolecules</u>: <u>The Theory and Practice of Sequence Comparison</u>, Addison-Wesley, Reading, MA; and software packages from NCBI, NIH; and the University of Wisconsin Genetics Computer Group (GCG), Madison, WI; each of which is incorporated herein by reference. This changes when considering conservative substitutions as matches.

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Conservative substitutions typically include substitutions within the following groups: glycine, alanine; valine, isoleucine, leucine; aspartic acid, glutamic acid; asparagine, glutamine; serine, threonine; lysine, arginine; and phenylalanine, tyrosine. Homologous amino acid 5 sequences are intended to include natural allelic and interspecies variations in the cytokine sequence. Typical homologous proteins or peptides will have from 50-100% homology (if gaps can be introduced), to 60-100% homology (if conservative substitutions are included) with an amino 10 acid sequence segment of Table 1. Homology measures will be at least about 70%, generally at least 76%, more generally at least 81%, often at least 85%, more often at least 88%, typically at least 90%, more typically at least 15 92%, usually at least 94%, more usually at least 95%, preferably at least 96%, and more preferably at least 97%, and in particularly preferred embodiments, at least 98% or more. The degree of homology will vary with the length of the compared segments. Homologous proteins or peptides, 20 such as the allelic variants, will share most biological

As used herein, the term "biological activity" is used to describe, without limitation, effects on inflammatory responses, innate immunity, and/or morphogenic development by cytokine-like ligands. For example, these receptors should mediate phosphatase or phosphorylase activities, which activities are easily measured by standard procedures. See, e.g., Hardie, et al. (eds. 1995) The Protein Kinase FactBook vols. I and II, Academic Press, San Diego, CA; Hanks, et al. (1991) Meth. Enzymol. 200:38-62; Hunter, et al. (1992) Cell 70:375-388; Lewin (1990) Cell 61:743-752; Pines, et al. (1991) Cold Spring Harbor Symp. Quant. Biol. 56:449-463; and Parker, et al. (1993) Nature 363:736-738. The receptors, or portions thereof, may be useful as phosphate labeling enzymes to label general or specific substrates.

activities with the embodiments described in Table 1.

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The terms ligand, agonist, antagonist, and analog of, e.g., a DIRS1, include molecules that modulate the characteristic cellular responses to cytokine ligand proteins, as well as molecules possessing the more standard structural binding competition features of ligand-receptor interactions, e.g., where the receptor is a natural receptor or an antibody. The cellular responses likely are typically mediated through receptor tyrosine kinase pathways.

Also, a ligand is a molecule which serves either as a natural ligand to which said receptor, or an analog thereof, binds, or a molecule which is a functional analog of the natural ligand. The functional analog may be a ligand with structural modifications, or may be a wholly unrelated molecule which has a molecular shape which interacts with the appropriate ligand binding determinants. The ligands may serve as agonists or antagonists, see, e.g., Goodman, et al. (eds. 1990) Goodman & Gilman's: The Pharmacological Bases of Therapeutics, Pergamon Press, New York.

Rational drug design may also be based upon structural studies of the molecular shapes of a receptor or antibody and other effectors or ligands. See, e.g., Herz, et al. (1997) J. Recept. Signal Transduct. Res. 17:671-776; and 25 Chaiken, et al. (1996) Trends Biotechnol. 14:369-375. Effectors may be other proteins which mediate other functions in response to ligand binding, or other proteins which normally interact with the receptor. One means for determining which sites interact with specific other proteins is a physical structure determination, e.g., x-ray 30 crystallography or 2 dimensional NMR techniques. will provide guidance as to which amino acid residues form molecular contact regions. For a detailed description of protein structural determination, see, e.g., Blundell and Johnson (1976) Protein Crystallography, Academic Press, New 35

York, which is hereby incorporated herein by reference.

## II. Activities

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The cytokine receptor-like proteins will have a number of different biological activities, e.g., modulating cell proliferation, or in phosphate metabolism, being added to or removed from specific substrates, typically proteins. Such will generally result in modulation of an inflammatory function, other innate immunity response, or a morphological effect. The subunit will probably have a specific low affinity binding to the ligand.

The DIRS1 has the characteristic motifs of a receptor signaling through the JAK pathway. See, e.g., Ihle, et al. (1997) Stem Cells 15(suppl. 1):105-111; Silvennoinen, et al. (1997) APMIS 105:497-509; Levy (1997) Cytokine Growth Factor Review 8:81-90; Winston and Hunter (1996) Current Biol. 6:668-671; Barrett (1996) Baillieres Clin. Gastroenterol. 10:1-15; and Briscoe, et al. (1996) Philos. Trans. R. Soc. Lond. B. Biol. Sci. 351:167-171.

The biological activities of the cytokine receptor subunits will be related to addition or removal of phosphate moieties to substrates, typically in a specific manner, but occasionally in a non specific manner.

Substrates may be identified, or conditions for enzymatic activity may be assayed by standard methods, e.g., as described in Hardie, et al. (eds. 1995) The Protein Kinase FactBook vols. I and II, Academic Press, San Diego, CA; Hanks, et al. (1991) Meth. Enzymol. 200:38-62; Hunter, et al. (1992) Cell 70:375-388; Lewin (1990) Cell 61:743-752; Pines, et al. (1991) Cold Spring Harbor Symp. Quant. Biol. 56:449-463; and Parker, et al. (1993) Nature 363:736-738.

III. Nucleic Acids

This invention contemplates use of isolated nucleic acid or fragments, e.g., which encode these or closely related proteins, or fragments thereof, e.g., to encode a corresponding polypeptide, preferably one which is biologically active. In addition, this invention covers isolated or recombinant DNAs which encode such proteins or

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polypeptides having characteristic sequences of the DIRS1s. Typically, the nucleic acid is capable of hybridizing, under appropriate conditions, with a nucleic acid sequence segment shown in Table 1, but preferably not with a corresponding segment of other receptors described in Table

- 3. Said biologically active protein or polypeptide can be a full length protein, or fragment, and will typically have a segment of amino acid sequence highly homologous, e.g., exhibiting significant stretches of identity, to one shown
- in Table 1. Further, this invention covers the use of isolated or recombinant nucleic acid, or fragments thereof, which encode proteins having fragments which are equivalent to the DIRS1 proteins. The isolated nucleic acids can have the respective regulatory sequences in the 5' and 3'
- flanks, e.g., promoters, enhancers, poly-A addition signals, and others from the natural gene.

An "isolated" nucleic acid is a nucleic acid, e.g., an RNA, DNA, or a mixed polymer, which is substantially pure, e.g., separated from other components which naturally

- accompany a native sequence, such as ribosomes, polymerases, and flanking genomic sequences from the originating species. The term embraces a nucleic acid sequence which has been removed from its naturally occurring environment, and includes recombinant or cloned
- DNA isolates, which are thereby distinguishable from naturally occurring compositions, and chemically synthesized analogs or analogs biologically synthesized by heterologous systems. A substantially pure molecule includes isolated forms of the molecule, either completely or substantially pure.

An isolated nucleic acid will generally be a homogeneous composition of molecules, but will, in some embodiments, contain heterogeneity, preferably minor. This heterogeneity is typically found at the polymer ends or portions not critical to a desired biological function or

activity.

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A "recombinant" nucleic acid is typically defined either by its method of production or its structure. reference to its method of production, e.g., a product made by a process, the process is use of recombinant nucleic acid techniques, e.g., involving human intervention in the nucleotide sequence. Typically this intervention involves in vitro manipulation, although under certain circumstances it may involve more classical animal breeding techniques. Alternatively, it can be a nucleic acid made by generating 10 a sequence comprising fusion of two fragments which are not naturally contiguous to each other, but is meant to exclude products of nature, e.g., naturally occurring mutants as found in their natural state. Thus, for example, products made by transforming cells with an unnaturally occurring 15 vector is encompassed, as are nucleic acids comprising sequence derived using any synthetic oligonucleotide Such a process is often done to replace a codon process. with a redundant codon encoding the same or a conservative amino acid, while typically introducing or removing a 20 restriction enzyme sequence recognition site. Alternatively, the process is performed to join together nucleic acid segments of desired functions to generate a single genetic entity comprising a desired combination of functions not found in the commonly available natural forms, e.g., encoding a fusion protein. Restriction enzyme 25 recognition sites are often the target of such artificial manipulations, but other site specific targets, e.g., promoters, DNA replication sites, regulation sequences, control sequences, or other useful features may be 30 incorporated by design. A similar concept is intended for a recombinant, e.g., fusion, polypeptide. This will include a dimeric repeat. Specifically included are synthetic nucleic acids which, by genetic code redundancy, encode equivalent polypeptides to fragments of DIRS1 and 35 fusions of sequences from various different related molecules, e.g., other cytokine receptor family members.

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A "fragment" in a nucleic acid context is a contiguous segment of at least about 17 nucleotides, generally at least 21 nucleotides, more generally at least 25 nucleotides, ordinarily at least 30 nucleotides, more ordinarily at least 35 nucleotides, often at least 39 5 nucleotides, more often at least 45 nucleotides, typically at least 50 nucleotides, more typically at least 55 nucleotides, usually at least 60 nucleotides, more usually at least 66 nucleotides, preferably at least 72 10 nucleotides, more preferably at least 79 nucleotides, and in particularly preferred embodiments will be at least 85 or more nucleotides. Typically, fragments of different genetic sequences can be compared to one another over appropriate length stretches, particularly defined segments 15 such as the domains described below.

A nucleic acid which codes for a DIRS1 will be particularly useful to identify genes, mRNA, and cDNA species which code for itself or closely related proteins, as well as DNAs which code for polymorphic, allelic, or other genetic variants, e.g., from different individuals or related species. Preferred probes for such screens are those regions of the interleukin which are conserved between different polymorphic variants or which contain nucleotides which lack specificity, and will preferably be full length or nearly so. In other situations, polymorphic variant specific sequences will be more useful.

This invention further covers recombinant nucleic acid molecules and fragments having a nucleic acid sequence identical to or highly homologous to the isolated DNA set forth herein. In particular, the sequences will often be operably linked to DNA segments which control transcription, translation, and DNA replication. These additional segments typically assist in expression of the desired nucleic acid segment.

Homologous, or highly identical, nucleic acid sequences, when compared to one another, e.g., DIRS1 sequences, exhibit significant similarity. The standards

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for homology in nucleic acids are either measures for homology generally used in the art by sequence comparison or based upon hybridization conditions. Comparative hybridization conditions are described in greater detail below.

For sequence comparison, typically one sequence acts as a reference sequence, to which test sequences are compared. When using a sequence comparison algorithm, test and reference sequences are input into a computer, subsequence coordinates are designated, if necessary, and sequence algorithm program parameters are designated. The sequence comparison algorithm then calculates the percent sequence identity for the test sequence(s) relative to the reference sequence, based on the designated program parameters.

Optical alignment of sequences for comparison can be conducted, e.g., by the local homology algorithm of Smith and Waterman (1981) Adv. Appl. Math. 2:482, by the homology alignment algorithm of Needlman and Wunsch (1970) J. Mol. Biol. 48:443, by the search for similarity method of Pearson and Lipman (1988) Proc. Nat'l Acad. Sci. USA 85:2444, by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by visual inspection (see generally Ausubel et al., supra).

One example of a useful algorithm is PILEUP. PILEUP creates a multiple sequence alignment from a group of related sequences using progressive, pairwise alignments to show relationship and percent sequence identity. It also plots a tree or dendrogram showing the clustering relationships used to create the alignment. PILEUP uses a simplification of the progressive alignment method of Feng and Doolittle (1987) J. Mol. Evol. 35:351-360. The method used is similar to the method described by Higgins and Sharp (1989) CABIOS 5:151-153. The program can align up to 300 sequences, each of a maximum length of 5,000

nucleotides or amino acids. The multiple alignment procedure begins with the pairwise alignment of the two most similar sequences, producing a cluster of two aligned sequences. This cluster is then aligned to the next most related sequence or cluster of aligned sequences. clusters of sequences are aligned by a simple extension of the pairwise alignment of two individual sequences. final alignment is achieved by a series of progressive, pairwise alignments. The program is run by designating specific sequences and their amino acid or nucleotide coordinates for regions of sequence comparison and by designating the program parameters. For example, a reference sequence can be compared to other test sequences to determine the percent sequence identity relationship using the following parameters: default gap weight (3.00), default gap length weight (0.10), and weighted end gaps.

Another example of algorithm that is suitable for determining percent sequence identity and sequence similarity is the BLAST algorithm, which is described Altschul, et al. (1900) <u>J. Mol. Biol.</u> 215:403-410. Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information (http://www.npbi.nlm.nih.gov/). This algorithm involves first identifying high scoring sequence pairs (HSPs) by identifying short words of length W in the query sequence, which either match or satisfy some positivevalued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul, et al., These initial neighborhood word hits act as seeds for initiating searches to find\longer HSPs containing The word hits are then extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Extension of the word hits in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to kero or below, due to

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the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T, and X determine the sensitivity and speed of the alignment. The BLAST program uses as defaults a wordlength (W) of 11, the BLOSUM62 scoring matrix (see Henikoff and Henikoff (1989) Proc. Nat'l Acad. Sci. USA 89:10915) alignments (B) of 50, expectation (E) of 10, M=5 N=4, and a comparison of both strands.

In addition to calculating percent sequence identity, 10 the BLAST algorithm also performs a statistical analysis of the similarity between two sequences (see, e.g., Karlin and Altschul (1993) Proc. Nat'l Acad. Sci. USA 90:5873-5787). One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an 15 indication of the probability by which a match between two nucleotide or amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a reference sequence if the smallest sum probability in a comparison of the test nucleic acid to the reference 20 nucleic acid is less than about 0.1, more preferably less than about 0.01, and most preferably less than about 0.001.

A further indication that two nucleic acid sequences of polypeptides are substantially identical is that the polypeptide encoded by the first nucleic acid is immunologically cross reactive with the polypeptide encoded by the second nucleic acid, as described below. Thus, a polypeptide is typically substantially identical to a second polypeptide, for example, where the two peptides differ only by conservative substitutions. Another indication that two nucleic acid sequences are substantially identical is that the two molecules hybridize to each other under stringent conditions, as described below. Hybridization under stringent conditions should give a background of at least 2-fold over background, preferably at least 3-5 or more.

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Substantial identity in the nucleic acid sequence comparison context means either that the segments, or their complementary strands, when compared, are identical when optimally aligned, with appropriate nucleotide insertions or deletions, in at least about 60% of the nucleotides, 5 generally at least 66%, ordinarily at least 71%, often at least 76%, more often at least 80%, usually at least 84%, more usually at least 88%, typically at least 91%, more typically at least about 93%, preferably at least about 10 95%, more preferably at least about 96 to 98% or more, and in particular embodiments, as high at about 99% or more of the nucleotides, including, e.g., segments encoding structural domains such as the segments described below. Alternatively, substantial identity will exist when the 15 segments will hybridize under selective hybridization conditions, to a strand or its complement, typically using a sequence derived from Table 1. Typically, selective hybridization will occur when there is at least about 55% homology over a stretch of at least about 14 nucleotides, 20 more typically at least about 65%, preferably at least about 75%, and more preferably at least about 90%. Kanehisa (1984) Nucl. Acids Res. 12:203-213, which is incorporated herein by reference. The length of homology comparison, as described, may be over longer stretches, and in certain embodiments will be over a stretch of at least 25 about 17 nucleotides, generally at least about 20 nucleotides, ordinarily at least about 24 nucleotides, usually at least about 28 nucleotides, typically at least about 32 nucleotides, more typically at least about 40 nucleotides, preferably at least about 50 nucleotides, and 30 more preferably at least about 75 to 100 or more nucleotides. This includes, e.g., 125, 150, 175, 200, 225, 246, 273, and other lengths.

Stringent conditions, in referring to homology in the 35 hybridization context, will be stringent combined conditions of salt, temperature, organic solvents, and other parameters typically controlled in hybridization

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reactions. Stringent temperature conditions will usually include temperatures in excess of about 30°C, more usually in excess of about 37°C, typically in excess of about 45° C, more typically in excess of about 55°C, preferably in excess of about 65°C, and more preferably in excess of about 70° C. Stringent salt conditions will ordinarily be less than about 500 mM, usually less than about 400 mM, more usually less than about 300 mM, typically less than about 200 mM, preferably less than about 100 mM, and more preferably less than about 80 mM, even down to less than about 20 mM. However, the combination of parameters is much more important than the measure of any single parameter. See, e.g., Wetmur and Davidson (1968) J. Mol. Biol. 31:349-370, which is hereby incorporated herein by reference.

The isolated DNA can be readily modified by nucleotide substitutions, nucleotide deletions, nucleotide insertions, and inversions of nucleotide stretches. modifications result in novel DNA sequences which encode this protein or its derivatives. These modified sequences can be used to produce mutant proteins (muteins) or to enhance the expression of variant species. expression may involve gene amplification, increased transcription, increased translation, and other mechanisms. Such mutant DIRS1-like derivatives include predetermined or site-specific mutations of the protein or its fragments, including silent mutations using genetic code degeneracy. "Mutant DIRS1" as used herein encompasses a polypeptide otherwise falling within the homology definition of the DIRS1 as set forth above, but having an amino acid sequence which differs from that of other cytokine receptor-like proteins as found in nature, whether by way of deletion, substitution, or insertion. In particular, "site specific mutant DIRS1" encompasses a protein having substantial sequence identity with a protein of Table 1, and typically shares most of the biological activities or effects of the forms disclosed herein.

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Although site specific mutation sites are predetermined, mutants need not be site specific. Mammalian DIRS1 mutagenesis can be achieved by making amino acid insertions or deletions in the gene, coupled with expression. Substitutions, deletions, insertions, or many combinations may be generated to arrive at a final Insertions include amino- or carboxy- terminal fusions. Random mutagenesis can be conducted at a target codon and the expressed mammalian DIRS1 mutants can then be screened for the desired activity, providing some aspect of a structure-activity relationship. Methods for making substitution mutations at predetermined sites in DNA having a known sequence are well known in the art, e.g., by M13 primer mutagenesis. See also Sambrook, et al. (1989) and Ausubel, et al. (1987 and periodic Supplements).

The mutations in the DNA normally should not place coding sequences out of reading frames and preferably will not create complementary regions that could hybridize to produce secondary mRNA structure such as loops or hairpins.

The phosphoramidite method described by Beaucage and Carruthers (1981) Tetra. Letts. 22:1859-1862, will produce suitable synthetic DNA fragments. A double stranded fragment will often be obtained either by synthesizing the complementary strand and annealing the strand together under appropriate conditions or by adding the complementary strand using DNA polymerase with an appropriate primer sequence.

Polymerase chain reaction (PCR) techniques can often be applied in mutagenesis. Alternatively, mutagenesis primers are commonly used methods for generating defined mutations at predetermined sites. See, e.g., Innis, et al. (eds. 1990) PCR Protocols: A Guide to Methods and Applications Academic Press, San Diego, CA; and Dieffenbach and Dveksler (eds. 1995) PCR Primer: A Laboratory Manual Cold Spring Harbor Press, CSH, NY.

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#### Proteins, Peptides IV.

As described above, the present invention encompasses primate DIRS1, e.g., whose sequences are disclosed in Table 1, and described above. Allelic and other variants are also contemplated, including, e.g., fusion proteins combining portions of such sequences with others, including epitope tags and functional domains.

The present invention also provides recombinant proteins, e.g., heterologous fusion proteins using segments from these rodent proteins. A heterologous fusion protein is a fusion of proteins or segments which are naturally not normally fused in the same manner. Thus, the fusion product of a DIRS1 with another cytokine receptor is a continuous protein molecule having sequences fused in a typical peptide linkage, typically made as a single translation product and exhibiting properties, e.g., sequence or antigenicity, derived from each source peptide. A similar concept applies to heterologous nucleic acid sequences.

20 In addition, new constructs may be made from combining similar functional or structural domains from other related proteins, e.g., cytokine receptors or Toll-like receptors, including species variants. For example, ligand-binding or other segments may be "swapped" between different new 25 fusion polypeptides or fragments. See, e.g., Cunningham, et al. (1989) <u>Science</u> 243:1330-1336; and O'Dowd, et al. (1988) J. Biol. Chem. 263:15985-15992, each of which is incorporated herein by reference. Thus, new chimeric polypeptides exhibiting new combinations of specificities will result from the functional linkage of receptor-binding 30 specificities. For example, the ligand binding domains from other related receptor molecules may be added or substituted for other domains of this or related proteins. The resulting protein will often have hybrid function and 35 properties. For example, a fusion protein may include a targeting domain which may serve to provide sequestering of

the fusion protein to a particular subcellular organelle.

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Candidate fusion partners and sequences can be selected from various sequence data bases, e.g., GenBank; NCBI, NIH; and BCG, University of Wisconsin Biotechnology Computing Group, Madison, WI, which are each incorporated herein by reference.

The present invention particularly provides muteins which bind cytokine-like ligands, and/or which are affected in signal transduction. Structural alignment of human DIRS1 with other members of the cytokine receptor family show conserved features/residues. See Table 3. Alignment of the human DIRS1 sequence with other members of the cytokine receptor family indicates various structural and functionally shared features. See also, Bazan, et al. (1996) Nature 379:591; Lodi, et al. (1994) Science 263:1762-1766; Sayle and Milner-White (1995) TIBS 20:374-376; and Gronenberg, et al. (1991) Protein Engineering 4:263-269.

Substitutions with either mouse sequences or human sequences are particularly preferred. Conversely, conservative substitutions away from the ligand binding interaction regions will probably preserve most signaling activities; and conservative substitutions away from the intracellular domains will probably preserve most ligand binding properties.

"Derivatives" of the primate DIRS1 include amino acid sequence mutants, glycosylation variants, metabolic derivatives and covalent or aggregative conjugates with other chemical moieties. Covalent derivatives can be prepared by linkage of functionalities to groups which are found in the DIRS1 amino acid side chains or at the N- or C- termini, e.g., by means which are well known in the art. These derivatives can include, without limitation, aliphatic esters or amides of the carboxyl terminus, or of residues containing carboxyl side chains, O-acyl derivatives of hydroxyl group-containing residues, and N-acyl derivatives of the amino terminal amino acid or

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amino-group containing residues, e.g., lysine or arginine.

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Acyl groups are selected from the group of alkyl-moieties, including C3 to C18 normal alkyl, thereby forming alkanoyl aroyl species.

In particular, glycosylation alterations are included, e.g., made by modifying the glycosylation patterns of a polypeptide during its synthesis and processing, or in further processing steps. Particularly preferred means for accomplishing this are by exposing the polypeptide to glycosylating enzymes derived from cells which normally provide such processing, e.g., mammalian glycosylation Deglycosylation enzymes are also contemplated. enzymes. Also embraced are versions of the same primary amino acid sequence which have other minor modifications, including phosphorylated amino acid residues, e.g., phosphotyrosine, phosphoserine, or phosphothreonine.

A major group of derivatives are covalent conjugates of the receptors or fragments thereof with other proteins of polypeptides. These derivatives can be synthesized in recombinant culture such as N- or C-terminal fusions or by the use of agents known in the art for their usefulness in cross-linking proteins through reactive side groups. Preferred derivatization sites with cross-linking agents are at free amino groups, carbohydrate moieties, and cysteine residues.

Fusion polypeptides between the receptors and other 25 homologous or heterologous proteins are also provided. Homologous polypeptides may be fusions between different receptors, resulting in, for instance, a hybrid protein exhibiting binding specificity for multiple different cytokine ligands, or a receptor which may have broadened or 30 weakened specificity of substrate effect. Likewise, heterologous fusions may be constructed which would exhibit a combination of properties or activities of the derivative proteins. Typical examples are fusions of a reporter 35 polypeptide, e.g., luciferase, with a segment or domain of a receptor, e.g., a ligand-binding segment, so that the presence or location of a desired ligand may be easily

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determined. See, e.g., Dull, et al., U.S. Patent No. 4,859,609, which is hereby incorporated herein by reference. Other gene fusion partners include glutathione-S-transferase (GST), bacterial ß-galactosidase, trpE, Protein A, ß-lactamase, alpha amylase, alcohol dehydrogenase, and yeast alpha mating factor. See, e.g., Godowski, et al. (1988) <u>Science</u> 241:812-816.

The phosphoramidite method described by Beaucage and Carruthers (1981) Tetra. Letts. 22:1859-1862, will produce suitable synthetic DNA fragments. A double stranded fragment will often be obtained either by synthesizing the complementary strand and annealing the strand together under appropriate conditions or by adding the complementary strand using DNA polymerase with an appropriate primer sequence.

Such polypeptides may also have amino acid residues which have been chemically modified by phosphorylation, sulfonation, biotinylation, or the addition or removal of other moieties, particularly those which have molecular shapes similar to phosphate groups. In some embodiments, the modifications will be useful labeling reagents, or serve as purification targets, e.g., affinity ligands.

Fusion proteins will typically be made by either recombinant nucleic acid methods or by synthetic polypeptide methods. Techniques for nucleic acid manipulation and expression are described generally, for example, in Sambrook, et al. (1989) Molecular Cloning: A Laboratory Manual (2d ed.), Vols. 1-3, Cold Spring Harbor Laboratory, and Ausubel, et al. (eds. 1987 and periodic supplements) Current Protocols in Molecular Biology, Greene/Wiley, New York, which are each incorporated herein by reference. Techniques for synthesis of polypeptides are described, for example, in Merrifield (1963) J. Amer. Chem. Soc. 85:2149-2156; Merrifield (1986) Science 232: 341-347; and Atherton, et al. (1989) Solid Phase Peptide Synthesis: A Practical Approach, IRL Press, Oxford; each of which is incorporated herein by reference. See also Dawson, et al.

(1994) <u>Science</u> 266:776-779 for methods to make larger polypeptides.

This invention also contemplates the use of derivatives of a DIRS1 other than variations in amino acid sequence or glycosylation. Such derivatives may involve covalent or aggregative association with chemical moieties. These derivatives generally fall into three classes: (1) salts, (2) side chain and terminal residue covalent modifications, and (3) adsorption complexes, for example 10 with cell membranes. Such covalent or aggregative derivatives are useful as immunogens, as reagents in immunoassays, or in purification methods such as for affinity purification of a receptor or other binding molecule, e.g., an antibody. For example, a cytokine 15 ligand can be immobilized by covalent bonding to a solid support such as cyanogen bromide-activated Sepharose, by methods which are well known in the art, or adsorbed onto polyolefin surfaces, with or without glutaraldehyde cross-linking, for use in the assay or purification of an cytokine receptor, antibodies, or other similar molecules. 20 The ligand can also be labeled with a detectable group, for example radioiodinated by the chloramine T procedure, covalently bound to rare earth chelates, or conjugated to another fluorescent moiety for use in diagnostic assays.

25 An DIRS1 of this invention can be used as an immunogen for the production of antisera or antibodies specific, e.g., capable of distinguishing between other cytokine receptor family members, for the DIRS1 or various fragments The purified DIRS1 can be used to screen 30 monoclonal antibodies or antigen-binding fragments prepared by immunization with various forms of impure preparations containing the protein. Antibodies can typically be substituted with antigen binding fragments of natural antibodies, e.g., Fab, Fab2, Fv, etc. The purified DIRS1 35 can also be used as a reagent to detect antibodies generated in response to the presence of elevated levels of expression, or immunological disorders which lead to

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antibody production to the endogenous receptor. Additionally, DIRS1 fragments may also serve as immunogens to produce the antibodies of the present invention, as described immediately below. For example, this invention contemplates antibodies having binding affinity to or being raised against the amino acid sequences shown in Table 1, fragments thereof, or various homologous peptides. In particular, this invention contemplates antibodies having binding affinity to, or having been raised against, specific fragments which are predicted to be, or actually

specific fragments which are predicted to be, or actually are, exposed at the exterior protein surface of the native DIRS1.

The blocking of physiological response to the receptor ligands may result from the inhibition of binding of the ligand to the receptor, likely through competitive inhibition. Thus, in vitro assays of the present invention will often use antibodies or antigen binding segments of these antibodies, or fragments attached to solid phase substrates. These assays will also allow for the diagnostic determination of the effects of either ligand binding region mutations and modifications, or other mutations and modifications, e.g., which affect signaling or enzymatic function.

This invention also contemplates the use of

competitive drug screening assays, e.g., where neutralizing antibodies to the receptor or fragments compete with a test compound for binding to a ligand or other antibody. In this manner, the neutralizing antibodies or fragments can be used to detect the presence of a polypeptide which

shares one or more binding sites to a receptor and can also be used to occupy binding sites on a receptor that might otherwise bind a ligand.

## V. Making Nucleic Acids and Protein

DNA which encodes the protein or fragments thereof can be obtained by chemical synthesis, screening cDNA libraries, or by screening genomic libraries prepared from

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a wide variety of cell lines or tissue samples. Natural sequences can be isolated using standard methods and the sequences provided herein, e.g., in Table 1. Other species counterparts can be identified by hybridization techniques, or by various PCR techniques, combined with or by searching in sequence databases, e.g., GenBank.

This DNA can be expressed in a wide variety of host cells for the synthesis of a full-length receptor or fragments which can in turn, for example, be used to generate polyclonal or monoclonal antibodies; for binding studies; for construction and expression of modified ligand binding or kinase/phosphatase domains; and for structure/function studies. Variants or fragments can be expressed in host cells that are transformed or transfected with appropriate expression vectors. These molecules can be substantially free of protein or cellular contaminants, other than those derived from the recombinant host, and therefore are particularly useful in pharmaceutical compositions when combined with a pharmaceutically acceptable carrier and/or diluent. The protein, or portions thereof, may be expressed as fusions with other proteins.

Expression vectors are typically self-replicating DNA or RNA constructs containing the desired receptor gene or its fragments, usually operably linked to suitable genetic control elements that are recognized in a suitable host cell. These control elements are capable of effecting expression within a suitable host. The specific type of control elements necessary to effect expression will depend upon the eventual host cell used. Generally, the genetic control elements can include a prokaryotic promoter system or a eukaryotic promoter expression control system, and typically include a transcriptional promoter, an optional operator to control the onset of transcription, transcription enhancers to elevate the level of mRNA

transcription enhancers to elevate the level of mRNA expression, a sequence that encodes a suitable ribosome binding site, and sequences that terminate transcription

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and translation. Expression vectors also usually contain an origin of replication that allows the vector to replicate independently of the host cell.

The vectors of this invention include those which contain DNA which encodes a protein, as described, or a fragment thereof encoding a biologically active equivalent The DNA can be under the control of a viral polypeptide. promoter and can encode a selection marker. This invention further contemplates use of such expression vectors which are capable of expressing eukaryotic cDNA coding for such a protein in a prokaryotic or eukaryotic host, where the vector is compatible with the host and where the eukaryotic cDNA coding for the receptor is inserted into the vector such that growth of the host containing the vector expresses the cDNA in question. Usually, expression vectors are designed for stable replication in their host cells or for amplification to greatly increase the total number of copies of the desirable gene per cell. always necessary to require that an expression vector replicate in a host cell, e.g., it is possible to effect transient expression of the protein or its fragments in various hosts using vectors that do not contain a replication origin that is recognized by the host cell. is also possible to use vectors that cause integration of the protein encoding portion or its fragments into the host DNA by recombination.

Vectors, as used herein, comprise plasmids, viruses, bacteriophage, integratable DNA fragments, and other vehicles which enable the integration of DNA fragments into the genome of the host. Expression vectors are specialized vectors which contain genetic control elements that effect expression of operably linked genes. Plasmids are the most commonly used form of vector but all other forms of vectors which serve an equivalent function and which are, or

become, known in the art are suitable for use herein. See,
e.g., Pouwels, et al. (1985 and Supplements) Cloning
Vectors: A Laboratory Manual, Elsevier, N.Y., and

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Rodriguez, et al. (eds. 1988) <u>Vectors: A Survey of Molecular Cloning Vectors and Their Uses</u>, Buttersworth, Boston, which are incorporated herein by reference.

Transformed cells are cells, preferably mammalian, that have been transformed or transfected with receptor vectors constructed using recombinant DNA techniques. Transformed host cells usually express the desired protein or its fragments, but for purposes of cloning, amplifying, and manipulating its DNA, do not need to express the subject protein. This invention further contemplates culturing transformed cells in a nutrient medium, thus permitting the receptor to accumulate in the cell membrane. The protein can be recovered, either from the culture or, in certain instances, from the culture medium.

15 For purposes of this invention, nucleic sequences are operably linked when they are functionally related to each other. For example, DNA for a presequence or secretory leader is operably linked to a polypeptide if it is expressed as a preprotein or participates in directing the polypeptide to the cell membrane or in secretion of the polypeptide. A promoter is operably linked to a coding sequence if it controls the transcription of the polypeptide; a ribosome binding site is operably linked to a coding sequence if it is positioned to permit

translation. Usually, operably linked means contiguous and in reading frame, however, certain genetic elements such as repressor genes are not contiguously linked but still bind to operator sequences that in turn control expression.

Suitable host cells include prokaryotes, lower

eukaryotes, and higher eukaryotes. Prokaryotes include
both gram negative and gram positive organisms, e.g., E.

coli and B. subtilis. Lower eukaryotes include yeasts,
e.g., S. cerevisiae and Pichia, and species of the genus
Dictyostelium. Higher eukaryotes include established

tissue culture cell lines from animal cells, both of

non-mammalian origin, e.g., insect cells, and birds, and of mammalian origin, e.g., human, primates, and rodents.

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YCp-series).

Prokaryotic host-vector systems include a wide variety of vectors for many different species. As used herein, E. coli and its vectors will be used generically to include equivalent vectors used in other prokaryotes. A representative vector for amplifying DNA is pBR322 or many 5 of its derivatives. Vectors that can be used to express the receptor or its fragments include, but are not limited to, such vectors as those containing the lac promoter (pUC-series); trp promoter (pBR322-trp); Ipp promoter (the pIN-series); lambda-pP or pR promoters (pOTS); or hybrid 10 promoters such as ptac (pDR540). See Brosius, et al. (1988) "Expression Vectors Employing Lambda-, trp-, lac-, and Ipp-derived Promoters", in Vectors: A Survey of Molecular Cloning Vectors and Their Uses, (eds. Rodriguez and Denhardt), Buttersworth, Boston, Chapter 10, pp. 15 205-236, which is incorporated herein by reference.

Lower eukaryotes, e.g., yeasts and Dictyostelium, may be transformed with DIRS1 sequence containing vectors. purposes of this invention, the most common lower eukaryotic host is the baker's yeast, Saccharomyces cerevisiae. It will be used to generically represent lower eukaryotes although a number of other strains and species are also available. Yeast vectors typically consist of a replication origin (unless of the integrating type), a selection gene, a promoter, DNA encoding the receptor or its fragments, and sequences for translation termination, polyadenylation, and transcription termination. Suitable expression vectors for yeast include such constitutive promoters as 3-phosphoglycerate kinase and various other glycolytic enzyme gene promoters or such inducible promoters as the alcohol dehydrogenase 2 promoter or metallothionine promoter. Suitable vectors include derivatives of the following types: self-replicating low copy number (such as the YRp-series), self-replicating high copy number (such as the YEp-series); integrating types (such as the YIp-series), or mini-chromosomes (such as the

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Higher eukaryotic tissue culture cells are normally the preferred host cells for expression of the functionally active interleukin protein. In principle, many higher eukaryotic tissue culture cell lines are workable, e.g., insect baculovirus expression systems, whether from an invertebrate or vertebrate source. However, mammalian cells are preferred. Transformation or transfection and propagation of such cells has become a routine procedure. Examples of useful cell lines include HeLa cells, Chinese hamster ovary (CHO) cell lines, baby rat kidney (BRK) cell 10 lines, insect cell lines, bird cell lines, and monkey (COS) cell lines. Expression vectors for such cell lines usually include an origin of replication, a promoter, a translation initiation site, RNA splice sites (if genomic DNA is used), a polyadenylation site, and a transcription termination 15 These vectors also usually contain a selection gene or amplification gene. Suitable expression vectors may be plasmids, viruses, or retroviruses carrying promoters derived, e.g., from such sources as from adenovirus, SV40, parvoviruses, vaccinia virus, or cytomegalovirus. 20 Representative examples of suitable expression vectors include pCDNA1; pCD, see Okayama, et al. (1985) Mol. Cell Biol. 5:1136-1142; pMC1neo PolyA, see Thomas, et al. (1987) Cell 51:503-512; and a baculovirus vector such as pAC 373 25 or pAC 610.

For secreted proteins, an open reading frame usually encodes a polypeptide that consists of a mature or secreted product covalently linked at its N-terminus to a signal peptide. The signal peptide is cleaved prior to secretion of the mature, or active, polypeptide. The cleavage site can be predicted with a high degree of accuracy from empirical rules, e.g., von-Heijne (1986) Nucleic Acids Research 14:4683-4690 and Nielsen, et al. (1997) Protein Eng. 10:1-12, and the precise amino acid composition of the signal peptide often does not appear to be critical to its function, e.g., Randall, et al. (1989) Science 243:1156-1159; Kaiser et al. (1987) Science 235:312-317.

It will often be desired to express these polypeptides in a system which provides a specific or defined glycosylation pattern. In this case, the usual pattern will be that provided naturally by the expression system. However, the pattern will be modifiable by exposing the polypeptide, e.g., an unglycosylated form, to appropriate glycosylating proteins introduced into a heterologous expression system. For example, the receptor gene may be co-transformed with one or more genes encoding mammalian or other glycosylating enzymes. Using this approach, certain mammalian glycosylation patterns will be achievable in prokaryote or other cells.

The source of DIRS1 can be a eukaryotic or prokaryotic host expressing recombinant DIRS1, such as is described above. The source can also be a cell line such as mouse Swiss 3T3 fibroblasts, but other mammalian cell lines are also contemplated by this invention, with the preferred cell line being from the human species.

Now that the sequences are known, the primate DIRS1,

fragments, or derivatives thereof can be prepared by
conventional processes for synthesizing peptides. These
include processes such as are described in Stewart and
Young (1984) Solid Phase Peptide Synthesis, Pierce Chemical
Co., Rockford, IL; Bodanszky and Bodanszky (1984) The

- Practice of Peptide Synthesis, Springer-Verlag, New York; and Bodanszky (1984) The Principles of Peptide Synthesis, Springer-Verlag, New York; all of each which are incorporated herein by reference. For example, an azide process, an acid chloride process, an acid anhydride
- process, a mixed anhydride process, an active ester process (for example, p-nitrophenyl ester, N-hydroxysuccinimide ester, or cyanomethyl ester), a carbodiimidazole process, an oxidative-reductive process, or a
- dicyclohexylcarbodiimide (DCCD)/additive process can be used. Solid phase and solution phase syntheses are both applicable to the foregoing processes. Similar techniques can be used with partial DIRS1 sequences.

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The DIRS1 proteins, fragments, or derivatives are suitably prepared in accordance with the above processes as typically employed in peptide synthesis, generally either by a so-called stepwise process which comprises condensing an amino acid to the terminal amino acid, one by one in sequence, or by coupling peptide fragments to the terminal amino acid. Amino groups that are not being used in the coupling reaction typically must be protected to prevent coupling at an incorrect location.

If a solid phase synthesis is adopted, the C-terminal amino acid is bound to an insoluble carrier or support through its carboxyl group. The insoluble carrier is not particularly limited as long as it has a binding capability to a reactive carboxyl group. Examples of such insoluble carriers include halomethyl resins, such as chloromethyl resin or bromomethyl resin, hydroxymethyl resins, phenol resins, tert-alkyloxycarbonylhydrazidated resins, and the like.

An amino group-protected amino acid is bound in

sequence through condensation of its activated carboxyl
group and the reactive amino group of the previously formed
peptide or chain, to synthesize the peptide step by step.
After synthesizing the complete sequence, the peptide is
split off from the insoluble carrier to produce the

peptide. This solid-phase approach is generally described
by Merrifield, et al. (1963) in J. Am. Chem. Soc.
85:2149-2156, which is incorporated herein by reference.

The prepared protein and fragments thereof can be isolated and purified from the reaction mixture by means of peptide separation, for example, by extraction, precipitation, electrophoresis, various forms of chromatography, and the like. The receptors of this invention can be obtained in varying degrees of purity depending upon desired uses. Purification can be accomplished by use of the protein purification techniques disclosed herein, see below, or by the use of the antibodies herein described in methods of immunoabsorbant

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affinity chromatography. This immunoabsorbant affinity chromatography is carried out by first linking the antibodies to a solid support and then contacting the linked antibodies with solubilized lysates of appropriate cells, lysates of other cells expressing the receptor, or lysates or supernatants of cells producing the protein as a result of DNA techniques, see below.

Generally, the purified protein will be at least about 40% pure, ordinarily at least about 50% pure, usually at least about 60% pure, typically at least about 70% pure, more typically at least about 80% pure, preferable at least about 90% pure and more preferably at least about 95% pure, and in particular embodiments, 97%-99% or more. Purity will usually be on a weight basis, but can also be on a molar basis. Different assays will be applied as appropriate.

### VI. Antibodies

20 e.g., primate DIRS1 proteins and fragments thereof, both in naturally occurring native forms and in their recombinant forms, the difference being that antibodies to the active receptor are more likely to recognize epitopes which are only present in the native conformations. Denatured

25 antigen detection can also be useful in, e.g., Western analysis. Anti-idiotypic antibodies are also contemplated, which would be useful as agonists or antagonists of a natural receptor or an antibody.

Antibodies, including binding fragments and single

chain versions, against predetermined fragments of the
protein can be raised by immunization of animals with
conjugates of the fragments with immunogenic proteins.

Monoclonal antibodies are prepared from cells secreting the
desired antibody. These antibodies can be screened for
binding to normal or defective protein, or screened for
agonistic or antagonistic activity. These monoclonal
antibodies will usually bind with at least a KD of about 1

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mM, more usually at least about 300  $\mu\text{M},$  typically at least about 100 $\mu\text{M}$ , more typically at least about 30  $\mu\text{M}$ , preferably at least about 10  $\mu\text{M}$ , and more preferably at least about 3  $\mu M$  or better.

The antibodies, including antigen binding fragments, of this invention can have significant diagnostic or therapeutic value. They can be potent antagonists that bind to the receptor and inhibit binding to ligand or inhibit the ability of the receptor to elicit a biological response, e.g., act on its substrate. They also can be useful as non-neutralizing antibodies and can be coupled to toxins or radionuclides to bind producing cells, or cells localized to the source of the interleukin. Further, these antibodies can be conjugated to drugs or other therapeutic agents, either directly or indirectly by means of a linker.

The antibodies of this invention can also be useful in diagnostic applications. As capture or non-neutralizing antibodies, they might bind to the receptor without inhibiting ligand or substrate binding. As neutralizing antibodies, they can be useful in competitive binding They will also be useful in detecting or quantifying ligand. They may be used as reagents for Western blot analysis, or for immunoprecipitation or immunopurification of the respective protein.

Protein fragments may be joined to other materials, particularly polypeptides, as fused or covalently joined polypeptides to be used as immunogens. Mammalian cytokine receptors and fragments may be fused or covalently linked to a variety of immunogens, such as keyhole limpet

- hemocyanin, bovine serum albumin, tetanus toxoid, etc. 30 Microbiology, Hoeber Medical Division, Harper and Row, 1969; Landsteiner (1962) Specificity of Serological Reactions, Dover Publications, New York; and Williams, et al. (1967) Methods in Immunology and Immunochemistry, Vol.
- 1, Academic Press, New York; each of which are incorporated 35 herein by reference, for descriptions of methods of preparing polyclonal antisera. A typical method involves

hyperimmunization of an animal with an antigen. The blood of the animal is then collected shortly after the repeated immunizations and the gamma globulin is isolated.

In some instances, it is desirable to prepare monoclonal antibodies from various mammalian hosts, such as mice, rodents, primates, humans, etc. Description of techniques for preparing such monoclonal antibodies may be found in, e.g., Stites, et al. (eds.) Basic and Clinical Immunology (4th ed.), Lange Medical Publications, Los Altos, CA, and references cited therein; Harlow and Lane 10 (1988) Antibodies: A Laboratory Manual, CSH Press; Goding (1986) Monoclonal Antibodies: Principles and Practice (2d ed.) Academic Press, New York; and particularly in Kohler and Milstein (1975) in Nature 256: 495-497, which discusses one method of generating monoclonal antibodies. 15 these references is incorporated herein by reference. Summarized briefly, this method involves injecting an animal with an immunogen. The animal is then sacrificed and cells taken from its spleen, which are then fused with 20 The result is a hybrid cell or "hybridoma" myeloma cells. that is capable of reproducing in vitro. The population of hybridomas is then screened to isolate individual clones, each of which secrete a single antibody species to the

immunogen. In this manner, the individual antibody species
obtained are the products of immortalized and cloned single
B cells from the immune animal generated in response to a
specific site recognized on the immunogenic substance.

Other suitable techniques involve in vitro exposure of lymphocytes to the antigenic polypeptides or alternatively to selection of libraries of antibodies in phage or similar vectors. See, Huse, et al. (1989) "Generation of a Large Combinatorial Library of the Immunoglobulin Repertoire in Phage Lambda," Science 246:1275-1281; and Ward, et al. (1989) Nature 341:544-546, each of which is hereby

incorporated herein by reference. The polypeptides and antibodies of the present invention may be used with or without modification, including chimeric or humanized

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antibodies. Frequently, the polypeptides and antibodies will be labeled by joining, either covalently or noncovalently, a substance which provides for a detectable signal. A wide variety of labels and conjugation techniques are known and are reported extensively in both

the scientific and patent literature. Suitable labels include radionuclides, enzymes, substrates, cofactors, inhibitors, fluorescent moieties, chemiluminescent moieties, magnetic particles, and the like. Patents,

teaching the use of such labels include U.S. Patent Nos. 10 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149; and 4,366,241. Also, recombinant or chimeric immunoglobulins may be produced, see Cabilly, U.S. Patent No. 4,816,567; or made in transgenic mice, see, e.g.,

Mendez, et al. (1997) <u>Nature Genetics</u> 15:146-156. 15 references are incorporated herein by reference.

The antibodies of this invention can also be used for affinity chromatography in isolating the DIRS1 proteins or peptides. Columns can be prepared where the antibodies are linked to a solid support, e.g., particles, such as agarose, Sephadex, or the like, where a cell lysate may be passed through the column, the column washed, followed by increasing concentrations of a mild denaturant, whereby the purified protein will be released. The protein may be used to purify antibody. Conversely, the antibodies may be immunoselected or immunodepleted to provide binding compositions of defined specificities.

The antibodies may also be used to screen expression libraries for particular expression products. Usually the antibodies used in such a procedure will be labeled with a moiety allowing easy detection of presence of antigen by antibody binding.

Antibodies raised against a cytokine receptor will also be used to raise anti-idiotypic antibodies. will be useful in detecting or diagnosing various 35 immunological conditions related to expression of the protein or cells which express the protein. They also will

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be useful as agonists or antagonists of the ligand, which may be competitive inhibitors or substitutes for naturally occurring ligands.

A cytokine receptor protein that specifically binds to or that is specifically immunoreactive with an antibody generated against a defined immunogen, such as an immunogen consisting of the amino acid sequence of SEQ ID NO: 2, is typically determined in an immunoassay. The immunoassay typically uses a polyclonal antiserum which was raised, e.g., to a protein of SEQ ID NO: 2. This antiserum is selected to have low crossreactivity against other cytokine receptor family members, e.g., IL-12 receptor beta or gp130, preferably from the same species, and any such crossreactivity is removed by immunoabsorption prior to use in the immunoassay.

In order to produce antisera for use in an immunoassay, the protein, e.g., of SEQ ID NO: 2, is isolated as described herein. For example, recombinant protein may be produced in a mammalian cell line. An appropriate host, e.g., an inbred strain of mice such as Balb/c, is immunized with the selected protein, typically using a standard adjuvant, such as Freund's adjuvant, and a standard mouse immunization protocol (see Harlow and Lane, supra). Alternatively, a synthetic peptide derived from the sequences disclosed herein and conjugated to a carrier protein can be used an immunogen. Polyclonal sera are collected and titered against the immunogen protein in an immunoassay, e.g., a solid phase immunoassay with the immunogen immobilized on a solid support. Polyclonal antisera with a titer of  $10^4$  or greater are selected and tested for their cross reactivity against other cytokine receptor family members, e.g., IL-12 receptor beta and/or gp130, using a competitive binding immunoassay such as the one described in Harlow and Lane, supra, at pages 570-573. Preferably at least two cytokine receptor family members

Preferably at least two cytokine receptor family members are used in this determination. These cytokine receptor family members can be produced as recombinant proteins and

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isolated using standard molecular biology and protein chemistry techniques as described herein.

Immunoassays in the competitive binding format can be used for the crossreactivity determinations. For example, the protein of SEQ ID NO: 2 can be immobilized to a solid 5 support. Proteins added to the assay compete with the binding of the antisera to the immobilized antigen. ability of the above proteins to compete with the binding of the antisera to the immobilized protein is compared to the proteins of IL-12 receptor beta or gp130. The percent crossreactivity for the above proteins is calculated, using standard calculations. Those antisera with less than 10% crossreactivity with each of the proteins listed above are selected and pooled. The cross-reacting antibodies are then removed from the pooled antisera by immunoabsorption with the above-listed proteins.

The immunoabsorbed and pooled antisera are then used in a competitive binding immunoassay as described above to compare a second protein to the immunogen protein (e.g., the DIRS1 like protein of SEQ ID NO: 2). In order to make this comparison, the two proteins are each assayed at a wide range of concentrations and the amount of each protein required to inhibit 50% of the binding of the antisera to the immobilized protein is determined. If the amount of the second protein required is less than twice the amount of the protein of the selected protein or proteins that is required, then the second protein is said to specifically bind to an antibody generated to the immunogen.

It is understood that these cytokine receptor proteins 30 are members of a family of homologous proteins that comprise at least 6 so far identified genes. particular gene product, such as the DIRS1, the term refers not only to the amino acid sequences disclosed herein, but also to other proteins that are allelic, non-allelic, or species variants. 'It is also understood that the terms 35 include nonnatural mutations introduced by deliberate mutation using conventional recombinant technology such as

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single site mutation, or by excising short sections of DNA encoding the respective proteins, or by substituting new amino acids, or adding new amino acids. Such minor alterations typically will substantially maintain the immunoidentity of the original molecule and/or its 5 biological activity. Thus, these alterations include proteins that are specifically immunoreactive with a designated naturally occurring DIRS1 protein. biological properties of the altered proteins can be determined by expressing the protein in an appropriate cell 10 line and measuring the appropriate effect, e.g., upon transfected lymphocytes. Particular protein modifications considered minor would include conservative substitution of amino acids with similar chemical properties, as described above for the cytokine receptor family as a whole. aligning a protein optimally with the protein of the cytokine receptors and by using the conventional immunoassays described herein to determine immunoidentity, one can determine the protein compositions of the invention.

## Kits and quantitation

Both naturally occurring and recombinant forms of the cytokine receptor like molecules of this invention are particularly useful in kits and assay methods. 25 example, these methods would also be applied to screening for binding activity, e.g., ligands for these proteins. Several methods of automating assays have been developed in recent years so as to permit screening of tens of thousands 30 of compounds per year. See, e.g., a BIOMEK automated workstation, Beckman Instruments, Palo Alto, California, and Fodor, et al. (1991) <u>Science</u> 251:767-773, which is incorporated herein by reference. The latter describes means for testing binding by a plurality of defined polymers synthesized on a solid substrate. The development 35 of suitable assays to screen for a ligand or agonist/antagonist homologous proteins can be greatly

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facilitated by the availability of large amounts of purified, soluble cytokine receptors in an active state such as is provided by this invention.

Purified DIRS1 can be coated directly onto plates for use in the aforementioned ligand screening techniques. However, non-neutralizing antibodies to these proteins can be used as capture antibodies to immobilize the respective receptor on the solid phase, useful, e.g., in diagnostic uses.

This invention also contemplates use of DIRS1, 10 fragments thereof, peptides, and their fusion products in a variety of diagnostic kits and methods for detecting the presence of the protein or its ligand. Alternatively, or additionally, antibodies against the molecules may be incorporated into the kits and methods. 15 Typically the kit will have a compartment containing either a DIRS1 peptide or gene segment or a reagent which recognizes one or the Typically, recognition reagents, in the case of peptide, would be a receptor or antibody, or in the case of a gene segment, would usually be a hybridization probe.

A preferred kit for determining the concentration of DIRS1 in a sample would typically comprise a labeled compound, e.g., ligand or antibody, having known binding affinity for DIRS1, a source of DIRS1 (naturally occurring or recombinant) as a positive control, and a means for separating the bound from free labeled compound, for example a solid phase for immobilizing the DIRS1 in the test sample. Compartments containing reagents, and instructions, will normally be provided.

30 Antibodies, including antigen binding fragments, specific for mammalian DIRS1 or a peptide fragment, or receptor fragments are useful in diagnostic applications to · detect the presence of elevated levels of ligand and/or its fragments. Diagnostic assays may be homogeneous (without a separation step between free reagent and antibody-antigen 35 complex) or heterogeneous (with a separation step). Various commercial assays exist, such as radioimmunoassay

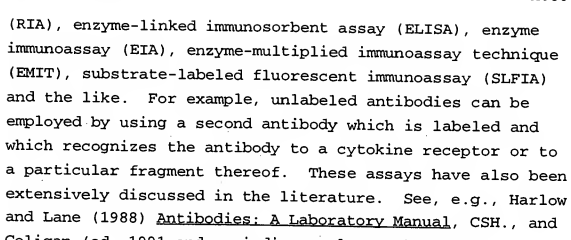
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10 Coligan (ed. 1991 and periodic supplements) <u>Current</u>

<u>Protocols In Immunology</u> Greene/Wiley, New York.

Anti-idiotypic antibodies may have similar use to

serve as agonists or antagonists of cytokine receptors. These should be useful as therapeutic reagents under appropriate circumstances.

Frequently, the reagents for diagnostic assays are supplied in kits, so as to optimize the sensitivity of the assay. For the subject invention, depending upon the nature of the assay, the protocol, and the label, either labeled or unlabeled antibody, or labeled ligand is This is usually in conjunction with other additives, such as buffers, stabilizers, materials necessary for signal production such as substrates for enzymes, and the like. Preferably, the kit will also contain instructions for proper use and disposal of the contents after use. Typically the kit has compartments for each useful reagent, and will contain instructions for proper use and disposal of reagents. Desirably, the reagents are provided as a dry lyophilized powder, where the reagents may be reconstituted in an aqueous medium having appropriate concentrations for performing the assay.

The aforementioned constituents of the diagnostic assays may be used without modification or may be modified in a variety of ways. For example, labeling may be achieved by covalently or non-covalently joining a moiety which directly or indirectly provides a detectable signal. In many of these assays, a test compound, cytokine

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receptor, or antibodies thereto can be labeled either directly or indirectly. Possibilities for direct labeling include label groups: radiolabels such as \$125\text{I}\$, enzymes (U.S. Pat. No. 3,645,090) such as peroxidase and alkaline phosphatase, and fluorescent labels (U.S. Pat. No. 3,940,475) capable of monitoring the change in fluorescence intensity, wavelength shift, or fluorescence polarization. Both of the patents are incorporated herein by reference. Possibilities for indirect labeling include biotinylation of one constituent followed by binding to avidin coupled to one of the above label groups.

There are also numerous methods of separating the bound from the free ligand, or alternatively the bound from the free test compound. The cytokine receptor can be immobilized on various matrixes followed by washing. Suitable matrices include plastic such as an ELISA plate, filters, and beads. Methods of immobilizing the receptor to a matrix include, without limitation, direct adhesion to plastic, use of a capture antibody, chemical coupling, and biotin-avidin. The last step in this approach involves the precipitation of antibody/antigen complex by any of several methods including those utilizing, e.g., an organic solvent such as polyethylene glycol or a salt such as ammonium sulfate. Other suitable separation techniques include, without limitation, the fluorescein antibody magnetizable particle method described in Rattle, et al. (1984) Clin. Chem. 30(9):1457-1461, and the double antibody magnetic particle separation as described in U.S. Pat. No. 4,659,678, each of which is incorporated herein by reference.

The methods for linking protein or fragments to various labels have been extensively reported in the literature and do not require detailed discussion here. Many of the techniques involve the use of activated carboxyl groups either through the use of carbodiimide or active esters to form peptide bonds, the formation of thioethers by reaction of a mercapto group with an

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activated halogen such as chloroacetyl, or an activated olefin such as maleimide, for linkage, or the like. Fusion proteins will also find use in these applications.

Another diagnostic aspect of this invention involves 5 use of oligonucleotide or polynucleotide sequences taken from the sequence of an cytokine receptor. These sequences can be used as probes for detecting levels of the respective cytokine receptor in patients suspected of having an immunological disorder. The preparation of both 10 RNA and DNA nucleotide sequences, the labeling of the sequences, and the preferred size of the sequences has received ample description and discussion in the literature. Normally an oligonucleotide probe should have at least about 14 nucleotides, usually at least about 18 15 nucleotides, and the polynucleotide probes may be up to several kilobases. Various labels may be employed, most commonly radionuclides, particularly <sup>32</sup>P. However, other techniques may also be employed, such as using biotin modified nucleotides for introduction into a 20 polynucleotide. The biotin then serves as the site for binding to avidin or antibodies, which may be labeled with a wide variety of labels, such as radionuclides, fluorescers, enzymes, or the like. Alternatively, antibodies may be employed which can recognize specific 25 duplexes, including DNA duplexes, RNA duplexes, DNA-RNA hybrid duplexes, or DNA-protein duplexes. The antibodies in turn may be labeled and the assay carried out where the duplex is bound to a surface, so that upon the formation of duplex on the surface, the presence of antibody bound to the duplex can be detected. The use of probes to the novel 30 anti-sense RNA may be carried out in conventional techniques such as nucleic acid hybridization, plus and minus screening, recombinational probing, hybrid released

35 This also includes amplification techniques such as polymerase chain reaction (PCR).

translation (HRT), and hybrid arrested translation (HART).

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Diagnostic kits which also test for the qualitative or quantitative presence of other markers are also contemplated. Diagnosis or prognosis may depend on the combination of multiple indications used as markers. Thus, kits may test for combinations of markers. See, e.g., Viallet, et al. (1989) Progress in Growth Factor Res. 1:89-97.

## VIII. Therapeutic Utility

This invention provides reagents with significant therapeutic value. See, e.g., Levitzki (1996) Curr. Opin. Cell Biol. 8:239-244. The cytokine receptors (naturally occurring or recombinant), fragments thereof, mutein receptors, and antibodies, along with compounds identified as having binding affinity to the receptors or antibodies, should be useful in the treatment of conditions exhibiting abnormal expression of the receptors of their ligands. Such abnormality will typically be manifested by immunological disorders. Additionally, this invention should provide therapeutic value in various diseases or disorders associated with abnormal expression or abnormal triggering of response to the ligand. For example, the IL-1 ligands have been suggested to be involved in morphologic development, e.g., dorso-ventral polarity determination, and immune responses, particularly the primitive innate responses. See, e.g., Sun, et al. (1991) Eur. J. Biochem. 196:247-254; and Hultmark (1994) Nature 367:116-117.

Recombinant cytokine receptors, muteins, agonist or antagonist antibodies thereto, or antibodies can be purified and then administered to a patient. reagents can be combined for therapeutic use with additional active ingredients, e.g., in conventional pharmaceutically acceptable carriers or diluents, along with physiologically innocuous stabilizers and excipients. These combinations can be sterile, e.g., filtered, and placed into dosage forms as by lyophilization in dosage

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vials or storage in stabilized aqueous preparations. This invention also contemplates use of antibodies or binding fragments thereof which are not complement binding.

Ligand screening using cytokine receptor or fragments thereof can be performed to identify molecules having binding affinity to the receptors. Subsequent biological assays can then be utilized to determine if a putative ligand can provide competitive binding, which can block intrinsic stimulating activity. Receptor fragments can be used as a blocker or antagonist in that it blocks the activity of ligand. Likewise, a compound having intrinsic stimulating activity can activate the receptor and is thus an agonist in that it simulates the activity of ligand, e.g., inducing signaling. This invention further contemplates the therapeutic use of antibodies to cytokine receptors as antagonists.

The quantities of reagents necessary for effective therapy will depend upon many different factors, including means of administration, target site, reagent physiological life, pharmacological life, physiological state of the patient, and other medicants administered. Thus, treatment dosages should be titrated to optimize safety and efficacy. Typically, dosages used in vitro may provide useful guidance in the amounts useful for in situ administration of these reagents. Animal testing of effective doses for treatment of particular disorders will provide further predictive indication of human dosage. considerations are described, e.g., in Gilman, et al. (eds. 1990) Goodman and Gilman's: The Pharmacological Bases of Therapeutics, 8th Ed., Pergamon Press; and Remington's Pharmaceutical Sciences, 17th ed. (1990), Mack Publishing Co., Easton, Penn.; each of which is hereby incorporated herein by reference. Methods for administration are discussed therein and below, e.g., for oral, intravenous, intraperitoneal, or intramuscular administration, transdermal diffusion, and others. Pharmaceutically

transdermal diffusion, and others. Pharmaceutically acceptable carriers will include water, saline, buffers,

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and other compounds described, e.g., in the Merck Index, Merck & Co., Rahway, New Jersey. Because of the likely high affinity binding, or turnover numbers, between a putative ligand and its receptors, low dosages of these reagents would be initially expected to be effective. And the signaling pathway suggests extremely low amounts of ligand may have effect. Thus, dosage ranges would ordinarily be expected to be in amounts lower than 1 mM concentrations, typically less than about 10 μM concentrations, usually less than about 100 nM, preferably less than about 1 fM (femtomolar), with an appropriate garrier

less than about 10 pM (picomolar), and most preferably less than about 1 fM (femtomolar), with an appropriate carrier. Slow release formulations, or slow release apparatus will often be utilized for continuous administration.

Cytokine receptors, fragments thereof, and antibodies or its fragments, antagonists, and agonists, may be administered directly to the host to be treated or, depending on the size of the compounds, it may be desirable to conjugate them to carrier proteins such as ovalbumin or serum albumin prior to their administration. Therapeutic formulations may be administered in many conventional dosage formulations. While it is possible for the active ingredient to be administered alone, it is preferable to present it as a pharmaceutical formulation. Formulations comprise at least one active ingredient, as defined above, together with one or more acceptable carriers thereof. Each carrier must be both pharmaceutically and physiologically acceptable in the sense of being compatible with the other ingredients and not injurious to the patient. Formulations include those suitable for oral, rectal, nasal, or parenteral (including subcutaneous, intramuscular, intravenous and intradermal) administration. The formulations may conveniently be presented in unit dosage form and may be prepared by methods well known in the art of pharmacy. See, e.g., Gilman, et al. (eds. 1990) Goodman and Gilman's: The Pharmacological Bases of

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Therapeutics, 8th Ed., Pergamon Press; and Remington's

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Pharmaceutical Sciences, 17th ed. (1990), Mack Publishing Co., Easton, Penn.; Avis, et al. (eds. 1993) Pharmaceutical Dosage Forms: Parenteral Medications Dekker, NY; Lieberman, et al. (eds. 1990) Pharmaceutical Dosage Forms: Tablets Dekker, NY; and Lieberman, et al. (eds. 1990) Pharmaceutical Dosage Forms: Disperse Systems Dekker, NY. The therapy of this invention may be combined with or used in association with other therapeutic agents, particularly agonists or antagonists of other cytokine receptor family members.

## IX. Screening

Drug screening using DIRS1 or fragments thereof can be performed to identify compounds having binding affinity to the receptor subunit, including isolation of associated components. Subsequent biological assays can then be utilized to determine if the compound has intrinsic stimulating activity and is therefore a blocker or antagonist in that it blocks the activity of the ligand. Likewise, a compound having intrinsic stimulating activity can activate the receptor and is thus an agonist in that it simulates the activity of a cytokine ligand. This invention further contemplates the therapeutic use of antibodies to the receptor as cytokine agonists or antagonists.

One method of drug screening utilizes eukaryotic or prokaryotic host cells which are stably transformed with recombinant DNA molecules expressing the DIRS1. Cells may be isolated which express a receptor in isolation from other functional receptors. Such cells, either in viable or fixed form, can be used for standard ligand/receptor binding assays. See also, Parce, et al. (1989) Science 246:243-247; and Owicki, et al. (1990) Proc. Nat'l Acad. Sci. USA 87:4007-4011, which describe sensitive methods to detect cellular responses. Competitive assays are particularly useful, where the cells (source of putative ligand) are contacted and incubated with a labeled receptor

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or antibody having known binding affinity to the ligand, such as <sup>125</sup>I-antibody, and a test sample whose binding affinity to the binding composition is being measured. The bound and free labeled binding compositions are then 5 separated to assess the degree of ligand binding. The amount of test compound bound is inversely proportional to the amount of labeled receptor binding to the known source. Any one of numerous techniques can be used to separate bound from free ligand to assess the degree of ligand This separation step could typically involve a 10 procedure such as adhesion to filters followed by washing, adhesion to plastic followed by washing, or centrifugation of the cell membranes. Viable cells could also be used to screen for the effects of drugs on cytokine mediated 15 functions, e.g., second messenger levels, i.e., Ca++; cell proliferation; inositol phosphate pool changes; and others. Some detection methods allow for elimination of a separation step, e.g., a proximity sensitive detection system. Calcium sensitive dyes will be useful for detecting Ca<sup>++</sup> levels, with a fluorimeter or a fluorescence 20 cell sorting apparatus.

## X. Ligands

The descriptions of the DIRS1 herein provide means to 25 identify ligands, as described above. Such ligand should bind specifically to the respective receptor with reasonably high affinity. Various constructs are made available which allow either labeling of the receptor to detect its ligand. For example, directly labeling cytokine 30 receptor, fusing onto it markers for secondary labeling, e.g., FLAG or other epitope tags, etc., will allow detection of receptor. This can be histological, as an affinity method for biochemical purification, or labeling or selection in an expression cloning approach. A twohybrid selection system may also be applied making 35 appropriate constructs with the available cytokine receptor

sequences. See, e.g., Fields and Song (1989) <u>Nature</u> 340:245-246.

Generally, descriptions of cytokine receptors will be analogously applicable to individual specific embodiments directed to DIRS1 reagents and compositions.

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The broad scope of this invention is best understood with reference to the following examples, which are not intended to limit the inventions to the specific embodiments.

#### **EXAMPLES**

#### I. General Methods

Some of the standard methods are described or 5 referenced, e.g., in Maniatis, et al. (1982) Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor Press; Sambrook, et al. (1989) Molecular Cloning: A Laboratory Manual, (2d ed.), vols. 1-3, CSH Press, NY; Ausubel, et al., Biology, Greene Publishing Associates, Brooklyn, NY; or Ausubel, et al. 10 (1987 and Supplements) <u>Current Protocols in Molecular</u> Biology, Greene/Wiley, New York. Methods for protein purification include such methods as ammonium sulfate precipitation, column chromatography, electrophoresis, 15 centrifugation, crystallization, and others. See, e.g., Ausubel, et al. (1987 and periodic supplements); Coligan, et al. (ed. 1996) and periodic supplements, <u>Current</u> Protocols In Protein Science Greene/Wiley, New York; Deutscher (1990) "Guide to Protein Purification" in Methods in Enzymology, vol. 182, and other volumes in this series; 20 and manufacturer's literature on use of protein purification products, e.g., Pharmacia, Piscataway, N.J., or Bio-Rad, Richmond, CA. Combination with recombinant techniques allow fusion to appropriate segments, e.g., to a 25 FLAG sequence or an equivalent which can be fused via a protease-removable sequence. See, e.g., Hochuli (1989) Chemische Industrie 12:69-70; Hochuli (1990) "Purification of Recombinant Proteins with Metal Chelate Absorbent" in Setlow (ed.) Genetic Engineering, Principle and Methods 12:87-98, Plenum Press, N.Y.; and Crowe, et al. (1992) 30 OIAexpress: The High Level Expression & Protein

Computer sequence analysis is performed, e.g., using available software programs, including those from the GCG (U. Wisconsin) and GenBank sources. Public sequence databases were also used, e.g., from GenBank and others.

Purification System QUIAGEN, Inc., Chatsworth, CA.

Many techniques applicable to IL-10 or IL-12 receptors may be applied to the DIRS1, as described, e.g., in USSN 08/110,683 (IL-10 receptor), which is incorporated herein by reference.

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## II. Computational Analysis

Human sequences related to cytokine receptors were identified from genomic sequence database using, e.g., the BLAST server (Altschul, et al. (1994) Nature Genet. 6:119-129). Standard analysis programs may be used to evaluate 10 structure, e.g., PHD (Rost and Sander (1994) Proteins 19:55-72) and DSC (King and Sternberg (1996) Protein Sci. 5:2298-2310). Standard comparison software includes, e.g., Altschul, et al. (1990) <u>J. Mol. Biol.</u> 215:403-10; Waterman 15 (1995) Introduction to Computational Biology: Maps, Sequences, and Genomes Chapman & Hall; Lander and Waterman (eds. 1995) Calculating the Secrets of Life: Applications of the Mathematical Sciences in Molecular Biology National Academy Press; and Speed and Waterman (eds. 1996) Genetic 20 Mapping and DNA Sequencing (Ima Volumes in Mathematics and Its Applications, Vol 81) Springer Verlag.

# III. Cloning of full-length DIRS cDNAs; Chromosomal localization

PCR primers derived from the DIRS sequences are used to probe a human cDNA library. Full length cDNAs for primate, rodent, or other species DIRS1 are cloned, e.g., by DNA hybridization screening of λgt10 phage. PCR reactions are conducted using T. aquaticus Taqplus DNA polymerase (Stratagene) under appropriate conditions.

Chromosome spreads are prepared. In situ hybridization is performed on chromosome preparations obtained from phytohemagglutinin-stimulated human lymphocytes cultured for 72 h. 5-bromodeoxyuridine was added for the final seven hours of culture (60  $\mu$ g/ml of medium), to ensure a posthybridization chromosomal banding of good quality.

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A PCR fragment, amplified with the help of primers, is cloned into an appropriate vector. The vector is labeled by nick-translation with <sup>3</sup>H. The radiolabeled probe is hybridized to metaphase spreads at final concentration of 200 ng/ml of hybridization solution as described in Mattei, et al. (1985) <u>Hum. Genet.</u> 69:327-331.

After coating with nuclear track emulsion (KODAK NTB<sub>2</sub>), slides are exposed. To avoid any slipping of silver grains during the banding procedure, chromosome spreads are first stained with buffered Giemsa solution and metaphase photographed. R-banding is then performed by the fluorochrome-photolysis-Giemsa (FPG) method and metaphases rephotographed before analysis. Alternatively, mapped sequence tags may be searched in a database.

Similar appropriate methods are used for other species.

## IV. Localization of DIRS1 or DIRS2 mRNA

Human multiple tissue (Cat# 1, 2) and cancer cell line 20 blots (Cat# 7757-1), containing approximately 2  $\mu g$  of poly(A) + RNA per lane, are purchased from Clontech (Palo Alto, CA). Probes are radiolabeled with  $[\alpha-32P]$  dATP, e.g., using the Amersham Rediprime random primer labeling kit (RPN1633). Prehybridization and hybridizations are performed at  $65^{\circ}$  C in 0.5 M Na<sub>2</sub>HPO<sub>4</sub>, 7% SDS, 0.5 M EDTA (pH 25 8.0). High stringency washes are conducted, e.g., at 65° C with two initial washes in 2 x SSC, 0.1% SDS for 40 min followed by a subsequent wash in 0.1 x SSC, 0.1% SDS for 20 Membranes are then exposed at -70° C to X-Ray film (Kodak) in the presence of intensifying screens. More 30 detailed studies by cDNA library Southerns are performed with selected human DIRS1 clones to examine their expression in hemopoietic or other cell subsets.

Alternatively, two appropriate primers are selected from Table 1 or 2. RT-PCR is used on an appropriate mRNA sample selected for the presence of message to produce a cDNA, e.g., a sample which expresses the gene.

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Full length clones may be isolated by hybridization of cDNA libraries from appropriate tissues pre-selected by PCR signal. Northern blots can be performed.

Message for genes encoding DIRS1 will be assayed by appropriate technology, e.g., PCR, immunoassay, hybridization, or otherwise. Tissue and organ cDNA preparations are available, e.g., from Clontech, Mountain View, CA. Identification of sources of natural expression are useful, as described. And the identification of functional receptor subunit pairings will allow for prediction of what cells express the combination of receptor subunits which will result in a physiological responsiveness to each of the cytokine ligands.

For mouse distribution, e.g., Southern Analysis can be 15 performed: DNA (5 µg) from a primary amplified cDNA library was digested with appropriate restriction enzymes to release the inserts, run on a 1% agarose gel and transferred to a nylon membrane (Schleicher and Schuell, Keene, NH).

Samples for mouse mRNA isolation may include: resting mouse fibroblastic L cell line (C200); Braf:ER (Braf fusion 20 to estrogen receptor) transfected cells, control (C201); T cells, TH1 polarized (Mell4 bright, CD4+ cells from spleen, polarized for 7 days with IFN-γ and anti IL-4; T200); T cells, TH2 polarized (Mell4 bright, CD4+ cells from spleen, polarized for 7 days with IL-4 and anti-IFN- $\gamma$ ; T201); T 25 cells, highly TH1 polarized (see Openshaw, et al. (1995) J. Exp. Med. 182:1357-1367; activated with anti-CD3 for 2, 6, 16 h pooled; T202); T cells, highly TH2 polarized (see Openshaw, et al. (1995) J. Exp. Med. 182:1357-1367; activated with anti-CD3 for 2, 6, 16 h pooled; T203); CD44-30 CD25+ pre T cells, sorted from thymus (T204); TH1 T cell clone D1.1, resting for 3 weeks after last stimulation with antigen (T205); TH1 T cell clone D1.1, 10  $\mu g/ml$  ConA stimulated 15 h (T206); TH2 T cell clone CDC35, resting for 3 weeks after last stimulation with antigen (T207); TH2 T 35 cell clone CDC35, 10  $\mu$ g/ml ConA stimulated 15 h (T208);

Mel14+ naive T cells from spleen, resting (T209); Mel14+ T

cells, polarized to Thl with IFN-y/IL-12/anti-IL-4 for 6, 12, 24 h pooled (T210); Mell4+ T cells, polarized to Th2 with IL-4/anti-IFN-γ for 6, 13, 24 h pooled (T211); unstimulated mature B cell leukemia cell line A20 (B200); unstimulated B cell line CH12 (B201); unstimulated large B 5 cells from spleen (B202); B cells from total spleen, LPS activated (B203); metrizamide enriched dendritic cells from spleen, resting (D200); dendritic cells from bone marrow, resting (D201); monocyte cell line RAW 264.7 activated with LPS 4 h (M200); bone-marrow macrophages derived with GM and 10 M-CSF (M201); macrophage cell line J774, resting (M202); macrophage cell line J774 + LPS + anti-IL-10 at 0.5, 1, 3, 6, 12 h pooled (M203); macrophage cell line J774 + LPS + IL-10 at 0.5, 1, 3, 5, 12 h pooled(M204); aerosol 15 challenged mouse lung tissue, Th2 primers, aerosol OVA challenge 7, 14, 23 h pooled (see Garlisi, et al. (1995) Clinical Immunology and Immunopathology 75:75-83; X206); Nippostrongulus-infected lung tissue (see Coffman, et al. (1989) Science 245:308-310; X200); total adult lung, normal (O200); total lung, rag-1 (see Schwarz, et al. (1993) 20 Immunodeficiency 4:249-252; O205); IL-10 K.O. spleen (see Kuhn, et al. (1991) Cell 75:263-274; X201); total adult spleen, normal (O201); total spleen, rag-1 (O207); IL-10 K.O. Peyer's patches (0202); total Peyer's patches, normal 25 (O210); IL-10 K.O. mesenteric lymph nodes (X203); total mesenteric lymph nodes, normal (O211); IL-10 K.O. colon (X203); total colon, normal (O212); NOD mouse pancreas (see Makino, et al. (1980) <u>Jikken Dobutsu</u> 29:1-13; X205); total thymus, rag-1 (0208); total kidney, rag-1 (0209); total 30 heart, rag-1 (0202); total brain, rag-1 (0203); total

Samples for human mRNA isolation may include:
peripheral blood mononuclear cells (monocytes, T cells, NK
cells, granulocytes, B cells), resting (T100); peripheral
blood mononuclear cells, activated with anti-CD3 for 2, 6,
12 h pooled (T101); T cell, THO clone Mot 72, resting

testes, rag-1 (O204); total liver, rag-1 (O206); rat normal joint tissue (O300); and rat arthritic joint tissue (X300).



67 (T102); T cell, THO clone Mot 72, activated with anti-CD28 and anti-CD3 for 3, 6, 12 h pooled (T103); T cell, TH0 clone Mot 72, anergic treated with specific peptide for 2, 7, 12 h pooled (T104); T cell, TH1 clone HY06, resting 5 (T107); T cell, TH1 clone HY06, activated with anti-CD28 and anti-CD3 for 3, 6, 12 h pooled (T108); T cell, TH1 clone HY06, anergic treated with specific peptide for 2, 6, 12 h pooled (T109); T cel1, TH2 clone HY935, resting (T110); T cell, TH2 clone HY935, activated with anti-CD28 and anti-CD3 for 2, 7, 12 h pooled (T111); T cells CD4+CD45RO- T cells polarized 27 days in anti-CD28, IL-4, and anti IFN- $\gamma$ , TH2 polarized, activated with anti-CD3 and anti-CD28 4 h (T116); T cell tumor lines Jurkat and Hut78, resting (T117); T cell clones, pooled AD130.2, Tc783.12,

- Tc783.13, Tc783.58, Tc782.69, resting (T118); T cell random 15  $\gamma\delta$  T cell clones, resting (T119); Splenocytes, resting (B100); Splenocytes, activated with anti-CD40 and IL-4 (B101); B cell EBV lines pooled WT49, RSB, JY, CVIR, 721.221, RM3, HSY, resting (B102); B cell line JY,
- 20 activated with PMA and ionomycin for 1, 6 h pooled (B103); NK 20 clones pooled, resting (K100); NK 20 clones pooled, activated with PMA and ionomycin for 6 h (K101); NKL clone, derived from peripheral blood of LGL leukemia patient, IL-2 treated (K106); NK cytotoxic clone 640-A30-1, resting
- 25 (K107); hematopoietic precursor line TF1, activated with PMA and ionomycin for 1, 6 h pooled (C100); U937 premonocytic line, resting (M100); U937 premonocytic line, activated with PMA and ionomycin for 1, 6 h pooled (M101); elutriated monocytes, activated with LPS, IFNY, anti-IL-10
- 30 for 1, 2, 6, 12, 24 h pooled (M102); elutriated monocytes, activated with LPS, IFNY, IL-10 for 1, 2, 6, 12, 24 h pooled (M103); elutriated monocytes, activated with LPS, IFNY, anti-IL-10 for 4, 16 h pooled (M106); elutriated monocytes, activated with LPS, IFNy, IL-10 for 4, 16 h pooled (M107);
- 35 elutriated monocytes, activated LPS for 1 h (M108); elutriated monocytes, activated LPS for 6 h (M109); DC 70% CD1a+, from CD34+ GM-CSF, TNFα 12 days, resting (D101); DC



70% CD1a+, from CD34+ GM-CSF, TNF $\alpha$  12 days, activated with PMA and ionomycin for 1 hr (D102); DC 70% CD1a+, from CD34+ GM-CSF,  $TNF\alpha$  12 days, activated with PMA and ionomycin for 6 hr (D103); DC 95% CD1a+, from CD34+ GM-CSF, TNF $\alpha$  12 days 5 FACS sorted, activated with PMA and ionomycin for 1, 6 h pooled (D104); DC 95% CD14+, ex CD34+ GM-CSF, TNF $\alpha$  12 days FACS sorted, activated with PMA and ionomycin 1. 6 hr pooled (D105); DC CD1a+ CD86+, from CD34+ GM-CSF, TNF $\alpha$  12 days FACS sorted, activated with PMA and ionomycin for 1, 6 10 h pooled (D106); DC from monocytes GM-CSF, IL-4 5 days, resting (D107); DC from monocytes GM-CSF, IL-4 5 days, resting (D108); DC from monocytes GM-CSF, IL-4 5 days, activated LPS 4, 16 h pooled (D109); DC from monocytes GM-CSF, IL-4 5 days, activated TNFα, monocyte supe for 4, 16 h 15 pooled (D110); leiomyoma L11 benign tumor (X101); normal myometrium M5 (O115); malignant leiomyosarcoma GS1 (X103); lung fibroblast sarcoma line MRC5, activated with PMA and ionomycin for 1, 6 h pooled (C101); kidney epithelial carcinoma cell line CHA, activated with PMA and ionomycin 20 for 1, 6 h pooled (C102); kidney fetal 28 wk male (O100); lung fetal 28 wk male (0101); liver fetal 28 wk male (O102); heart fetal 28 wk male (O103); brain fetal 28 wk male (0104); gallbladder fetal 28 wk male (0106); small intestine fetal 28 wk male (0107); adipose tissue fetal 28 25 wk male (0108); ovary fetal 25 wk female (0109); uterus fetal 25 wk female (O110); testes fetal 28 wk male (O111); spleen fetal 28 wk male (0112); adult placenta 28 wk (0113); and tonsil inflamed, from 12 year old (X100).

With a cDNA Southern, the human DIRS1 was found in 30 LPS activated dendritic cells ("DC LPS"); monokine activated dendritic cells ("DC mix"); normal skin; Psoriasis skin; inflamed tonsil; fetal liver; fetal small intestine; fetal ovary; resting "70% dendritic cells"; 6 hr activated 70% dendritic cells; and LPS activated monocytes.

35 A signal was also detected in normal monkey lung and Ascaris-challenged monkey lung (24 h), which indicates

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cross species hybridization. The following libraries had weaker expression of DIRS1: smoker lung pool; fetal spleen CD4+ T cells (TH2 polarized); gamma delta T cells; activated splenocytes; and B cells.

HOFNy28 (DIRS2) is expressed in U937 (a premonocytic cell line) cells, both resting and activated; activated A549 cells (epithelial cells, IL-1 $\beta$  activated); fetal uterus; fetal testes; and fetal spleen. This data is from PCR on these cDNA libraries followed by Southern hybridization.

Similar samples may isolated in other species for evaluation.

- V. Cloning of species counterparts of DIRS1 or DIRS2

  Various strategies are used to obtain species
  counterparts of, e.g., the DIRS1, preferably from other
  primates or rodents. One method is by cross hybridization
  using closely related species DNA probes. It may be useful
  to go into evolutionarily similar species as intermediate

  20 steps. Another method is by using specific PCR primers
  based on the identification of blocks of similarity or
  difference between genes, e.g., areas of highly conserved
  or nonconserved polypeptide or nucleotide sequence.
  Database sequence searches may also identify species

  25 counterparts.
- VI. Production of mammalian DIRS1 or DIRS2 protein
  An appropriate, e.g., GST, fusion construct is
  engineered for expression, e.g., in E. coli. For example,
  a mouse IGIF pGex plasmid is constructed and transformed
  into E. coli. Freshly transformed cells are grown, e.g.,
  in LB medium containing 50 µg/ml ampicillin and induced
  with IPTG (Sigma, St. Louis, MO). After overnight
  induction, the bacteria are harvested and the pellets
  containing the DIRS1 protein are isolated. The pellets are
  homogenized, e.g., in TE buffer (50 mM Tris-base pH 8.0, 10
  mM EDTA and 2 mM pefabloc) in 2 liters. This material is

passed through a microfluidizer (Microfluidics, Newton, MA) three times. The fluidized supernatant is spun down on a Sorvall GS-3 rotor for 1 h at 13,000 rpm. The resulting supernatant containing the cytokine receptor protein is filtered and passed over a glutathione-SEPHAROSE column equilibrated in 50 mM Tris-base pH 8.0. The fractions containing the DIRS1-GST fusion protein are pooled and cleaved, e.g., with thrombin (Enzyme Research Laboratories, Inc., South Bend, IN). The cleaved pool is then passed over a O-SEPHAROSE column equilibrated in 50 mM Tris-base. Fractions containing DIRS1 are pooled and diluted in cold distilled H2O, to lower the conductivity, and passed back over a fresh Q-Sepharose column, alone or in succession with an immunoaffinity antibody column. Fractions containing the DIRS1 protein are pooled, aliquoted, and stored in the -70° C freezer.

Comparison of the CD spectrum with cytokine receptor protein may suggest that the protein is correctly folded. See Hazuda, et al. (1969) J. Biol. Chem. 264:1689-1693.

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Determining physiological forms of receptors The cellular forms of receptors for ligands can be tested with the various ligands and receptor subunits provided, e.g., IL-10 kalated sequences. In particular, multiple cytokine receptor like ligands have been identified, see, e.g., US\$N 60/027,368, 08/934,959, and

08/842,659, which are incomporated herein by reference...

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Cotransformation of the DIRS1 with putative other receptor subunit genes may be performed. In particular, the DSRS1 is suggested to be a second receptor subunit needed for functional receptor signaling. Such cells may be used to screen putative cytokine ligands, such as the DIL-30, for signaling. A cell proliferation assay may be used.

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In addition, it has been known that many cytokine receptors function as heterodimers. The IL-1 $\alpha$  and IL-1 $\beta$ ligands bind an IL-1R1 as the primary receptor and this

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complex then forms a high affinity receptor complex with the IL-1R3. As indicated above, the sequence similarity to IL-12 receptor subunits suggests functional similarity of the functional receptor, e.g., a soluble alpha subunit, and transmembrane beta subunit.

These subunit combinations can be tested now with the provided reagents. In particular, appropriate constructs can be made for transformation or transfection of subunits into cells. Constructs for the alpha chains, e.g., DSRS1 forms, can be made. Likewise for the beta subunit DIRS1. Combinatorial transfections of transformations can make cells expressing defined subunits, which can be tested for response to the predicted ligands. Appropriate cell types can be used, e.g., 293 T cells, with, e.g., an NFKb reporter construct.

Biological assays will generally be directed to the ligand binding feature of the protein or to the kinase/phosphatase activity of the receptor. The activity will typically be reversible, as are many other enzyme reactions, and may mediate phosphatase or phosphorylase activities, which activities are easily measured by standard procedures. See, e.g., Hardie, et al. (eds. 1995) The Protein Kinase FactBook vols. I and II, Academic Press, San Diego, CA; Hanks, et al. (1991) Meth. Enzymol. 200:38-62; Hunter, et al. (1992) Cell 70:375-388; Lewin (1990) Cell 61:743-752; Pines, et al. (1991) Cold Spring Harbor Symp. Quant. Biol. 56:449-463; and Parker, et al. (1993) Nature 363:736-738.

The family of cytokines contains molecules which are important mediators of hematopoiesis or inflammatory disease. See, e.g., Thomson (ed. 1994) The Cytokine Handbook Academic Press, San Diego; and Dinarello (1996) Blood 87:2095-2147.

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VIII. Antibodies specific for DIRS1 or DIRS2

Inbred Balb/c mice are immunized intraperitoneally with recombinant forms of the protein, e.g., purified DIRS1 or stable transfected NIH-3T3 cells. Animals are boosted at appropriate time points with protein, with or without additional adjuvant, to further stimulate antibody production. Serum is collected, or hybridomas produced with harvested spleens.

Alternatively, Balb/c mice are immunized with cells 10 transformed with the gene or fragments thereof, either endogenous or exogenous cells, or with isolated membranes enriched for expression of the antigen. Serum is collected at the appropriate time, typically after numerous further administrations. Various gene therapy techniques may be 15 useful, e.g., in producing protein in situ, for generating an immune response. Serum may be immunoselected or depleted to prepare substantially purified antibodies of defined specificity and high affinity. Preparations which specifically bind particular segments or defined epitopes 20 may be made.

Monoclonal antibodies may be made. For example, splenocytes are fused with an appropriate fusion partner and hybridomas are selected in growth medium by standard procedures. Hybridoma supernatants are screened for the presence of antibodies which bind to the DIRS1, e.g., by ELISA or other assay. Antibodies which specifically recognize specific DIRS1 embodiments may also be selected or prepared.

In another method, synthetic peptides or purified

protein are presented to an immune system to generate monoclonal or polyclonal antibodies. See, e.g., Coligan (ed. 1991) Current Protocols in Immunology Wiley/Greene; and Harlow and Lane (1989) Antibodies: A Laboratory Manual Cold Spring Harbor Press. In appropriate situations, the binding reagent is either labeled as described above, e.g., fluorescence or otherwise, or immobilized to a substrate for panning methods. Nucleic acids may also be introduced

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into cells in an animal to produce the antigen, which serves to elicit an immune response. See, e.g., Wang, et al. (1993) Proc. Nat'l. Acad. Sci. 90:4156-4160; Barry, et al. (1994) BioTechniques 16:616-619; and Xiang, et al. (1995) Immunity 2: 129-135.

Moreover, antibodies which may be useful to determine the combination of the DIRS1 with a functional alpha subunit may be generated. Thus, e.g., epitopes characteristic of a particular functional alpha/beta combination may be identified with appropriate antibodies.

IX. Production of fusion proteins with DIRS1 or DIRS2

Various fusion constructs are made with DIRS1 or

DIRS2. A portion of the appropriate gene is fused to an
epitope tag, e.g., a FLAG tag, or to a two hybrid system
construct. See, e.g., Fields and Song (1989) Nature
340:245-246.

The epitope tag may be used in an expression cloning procedure with detection with anti-FLAG antibodies to detect a binding partner, e.g., ligand for the respective cytokine receptor. The two hybrid system may also be used to isolate proteins which specifically bind to DIRS1.

## X. Structure activity relationship

Information on the criticality of particular residues is determined using standard procedures and analysis. Standard mutagenesis analysis is performed, e.g., by generating many different variants at determined positions, e.g., at the positions identified above, and evaluating

biological activities of the variants. This may be performed to the extent of determining positions which modify activity, or to focus on specific positions to determine the residues which can be substituted to either retain, block, or modulate biological activity.

Alternatively, analysis of natural variants can indicate what positions tolerate natural mutations. This may result from populational analysis of variation among

PARHAM, et al.

individuals, or across strains or species. Samples from selected individuals are analyzed, e.g., by PCR analysis and sequencing. This allows evaluation of population polymorphisms.

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#### Isolation of a ligand for DIRS1 or DIRS2 XI.

A cytokine receptor can be used as a specific binding reagent to identify its binding partner, by taking advantage of its specificity of binding, much like an antibody would be used. Typically, the binding receptor is a heterodimer of receptor subunits. A binding reagent is either labeled as described above, e.g., fluorescence or otherwise, or immobilized to a substrate for panning methods.

15 The binding composition is used to screen an expression library made from a cell line which expresses a binding partner, i.e., ligand, preferably membrane associated. Standard staining techniques are used to detect or sort surface expressed ligand, or surface 20 expressing transformed cells are screened by panning. Screening of intracellular expression is performed by various staining or immunofluorescence procedures. also McMahan, et al. (1991) EMBO J. 10:2821-2832.

For example, on day 0, precoat 2-chamber permanox 25 slides with 1 ml per chamber of fibronectin, 10 ng/ml in PBS, for 30 min at room temperature. Rinse once with PBS. Then plate COS cells at 2-3  $\times$   $10^5$  cells per chamber in 1.5 ml of growth media. Incubate overnight at 37° C.

On day 1 for each sample, prepare 0.5 ml of a solution of 66  $\mu$ g/ml DEAE-dextran, 66  $\mu$ M chloroquine, and 4  $\mu$ g DNA in serum free DME. For each set, a positive control is prepared, e.g., of DIRS1-FLAG cDNA at 1 and 1/200 dilution, and a negative mock. Rinse cells with serum free DME. the DNA solution and incubate 5 hr at 37° C. Remove the medium and add 0.5 ml 10% DMSO in DME for 2.5 min. and wash once with DME. Add 1.5 ml growth medium and incubate overnight.

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On day 2, change the medium. On days 3 or 4, the cells are fixed and stained. Rinse the cells twice with Hank's Buffered Saline Solution (HBSS) and fix in 4% paraformaldehyde (PFA)/glucose for 5 min. Wash 3X with The slides may be stored at -80° C after all liquid 5 is removed. For each chamber, 0.5 ml incubations are performed as follows. Add HBSS/saponin (0.1%) with 32  $\mu$ l/ml of 1 M NaN3 for 20 min. Cells are then washed with HBSS/saponin 1X. Add appropriate DIRS1 or DIRS1/antibody complex to cells and incubate for 30 min. Wash cells twice 10 with HBSS/saponin. If appropriate, add first antibody for 30 min. Add second antibody, e.g., Vector anti-mouse antibody, at 1/200 dilution, and incubate for 30 min. Prepare ELISA solution, e.g., Vector Elite ABC horseradish peroxidase solution, and preincubate for 30 min. Use, 15 e.g., 1 drop of solution A (avidin) and 1 drop solution B (biotin) per 2.5 ml HBSS/saponin. Wash cells twice with HBSS/saponin. Add ABC HRP solution and incubate for 30 min. Wash cells twice with HBSS, second wash for 2 min, which closes cells. Then add Vector diaminobenzoic acid 20 (DAB) for 5 to 10 min. Use 2 drops of buffer plus 4 drops DAB plus 2 drops of  $H_2O_2$  per 5 ml of glass distilled water. Carefully remove chamber and rinse slide in water. Air dry for a few minutes, then add 1 drop of Crystal Mount and a 25

Evaluate positive staining of pools and progressively subclone to isolation of single genes responsible for the binding.

cover slip. Bake for 5 min at 85-90° C.

Alternatively, receptor reagents are used to affinity purify or sort out cells expressing a putative ligand. See, e.g., Sambrook, et al. or Ausubel, et al.

Another strategy is to screen for a membrane bound receptor by panning. The receptor cDNA is constructed as described above. The ligand can be immobilized and used to immobilize expressing cells. Immobilization may be achieved by use of appropriate antibodies which recognize, e.g., a FLAG sequence of a DIRS1 fusion construct, or by

use of antibodies raised against the first antibodies. Recursive cycles of selection and amplification lead to enrichment of appropriate clones and eventual isolation of receptor expressing clones.

Phage expression libraries can be screened by mammalian DIRS1. Appropriate label techniques, e.g., anti-FLAG antibodies, will allow specific labeling of appropriate clones.

All citations herein are incorporated herein by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference.

Many modifications and variations of this invention can be

made without departing from its spirit and scope, as will be
apparent to those skilled in the art. The specific embodiments
described herein are offered by way of example only, and the
invention is to be limited by the terms of the appended claims,
along with the full scope of equivalents to which such claims
are entitled; and the invention is not to be limited by the
specific embodiments that have been presented herein by way of
example.

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